

METHOD FOR EXAMINING REACTIVITY
AND METHOD FOR DETECTING A COMPLEX

5 This application is a division of U.S. Patent Application
No. 09/942,662, filed August 31, 2001, now abandoned, which is
incorporated herein by reference.

BACKGROUND OF THE INVENTION

Field of the Invention

10 The present invention is directed to examining multiple
specimens at a time for multiple items, and provides a method
in which matrix substrates with biological samples having
different properties and origins bound thereto are prepared,
and on each matrix region, oligonucleotides having different
15 sequences, proteins or drugs are spotted in an array, whereby
multiple specimens are examined at a time for multiple items.

 The present invention also relates to a method in which,
by using an oligonucleotide having a known base sequence as a
detection probe to detect whether a complex is formed by
20 intermolecular bond with this oligonucleotide, detection is
made as to whether or not components having a capability of
bonding to the above -described detection probe are contained,
and to a detection substrate having the oligonucleotide as a
detection probe fixed on its surface, which is used
25 exclusively for this detecting method.

Related Background Art

 In identification of partial sequences included in the
base sequence of a nucleic acid molecule, detection of a
target nucleic acid contained in a sample originated from an
30 organism or identification of genus or species for various
bacteria based on the characteristics of the gene DNA of the

5 bacteria, a procedure may be used in which two or more probe
DNAs having known base sequences are used to detect whether or
not the nucleic acid molecule is a nucleic acid molecule
specifically binding to each probe DNA, namely making
hybridization with each probe DNA. As an effective approach
10 to performing speedily and accurately examination of the two
or more probe DNAs by the hybridization method, a procedure is
proposed in which a probe array of two or more probe DNAs
arranged regularly on a solid phase is used to detect at a
time whether or not the nucleic acid molecule is a nucleic
15 acid molecule specifically binding to each probe DNA.

Among common methods for producing such probe arrays, as
described in European Patent No. 373203 (EP 0373203 B1) for
example, methods are known in which predetermined nucleic acid
probes are synthesized in an array form on a solid phase, and
20 methods in which a plurality of nucleic acid probes
synthesized in advance is supplied in an array form on the
solid phase.

Prior technical documents disclosing the former methods
include, for example, U.S. Patent No. 5,405,783. Also, as one
25 example of the latter methods, a method in which cDNAs are
arranged in an array form on a solid phase using
micropipetting is disclosed in, for example, U.S. Patent No.
5,601,980 and "Science", Vol. 270, pp. 467, (1995).

The probe array that is prepared ~~with~~using these methods
30 may be an array ~~such that~~in which nucleic acid probes are
arranged on a solid phase at a high density of 10000 or more
probes per square inch. ~~Hybridization~~A hybridization reaction
with multiple probes ~~are~~is carried out at a time by dipping
this high-density probe array into a specimen solution, and in
35 so doing, the base sequence of genes is analyzed based on the
base sequence of nucleic acids ~~making hybridization~~that

5 hybridize. This method has an advantage in that probes are
arranged ~~in~~ at a high density on a substrate ~~of~~ with a small
area, thereby making it possible to conduct a multiple-item
examination at ~~a~~ the same time with a small amount of samples
to reduce the burden associated with sampling from the
10 subject.

As a method of preparing the high-density probe array for
the above -described application on the substrate by the DNA
synthesis process, a method in which a photolithography
technology is applied is disclosed in the aforesaid U.S.
15 Patent No. 5,405,783, but highly advanced equipment is
required for implementing this method, and the method is not
easy enough for anyone to use.

Also, in the case where the number of specimens is large
but the number of required examination items is not so large,
20 the integration degree of DNA probes on the probe array
corresponding to the number of examination items does not need
to be very high. Rather, there may be cases where it is
necessary to prepare a large number of probe arrays with a
small number of desired DNA probes fixed, using a simpler
25 method.

Actually, in the field of clinical examination, there are
not necessarily many cases where examinations for more than
10000 items are required. For example, in the case of group
health examination and the like, there may be cases where it
30 is more important to examine a large number of specimens with
a limited number of items. For examining a large number of
specimens in this way, a system is required such that presence
of diseases can be speedily examined through comparison with
standard samples with respect to each specimen.

35 In addition, the amount of a DNA specimen is generally
small as compared to that of an oligonucleotide capable of

5 being synthesized and used in the probe. For using it in a normal form in which the probe array substrate is dipped into the specimen solution for a hybridization reaction, the amount of specimen DNA allowing the substrate to be dipped sufficiently is required. Therefore, the size of the DNA
10 probe array substrate is limited depending on the amount of specimen DNA, and thus the array needs to be highly dense. Alternatively, as a result of diluting the specimen solution to ensure its volume for the size of the probe array substrate, the concentration of DNA in the specimen solution
15 is reduced, and a procedure is adopted ~~of~~for prolonging the reaction time to ~~make compensation~~compensate for the reduced concentration.

Also, since the amount of sampled specimens is limited inherently because the specimen is an extract from tissues,
20 and because it is subjected to pre-processing for making a specimen solution for use in the hybridization reaction, specifically, the extraction of nucleic acid, single-strand formation thereof, and process for labeling, the amount of finally obtained samples is very small. In order to ~~make~~
25 ~~compensation~~compensate for that, the sample is subjected to processing for amplification of the amount of DNA, such as amplification processing by PCR reaction, before it is used for examination and studies. However, there exists a disadvantage in that because ~~primers~~ separately prepared
30 primers are required for carrying out a PCR reaction, such processing can be applied only to specific genes of which primer sequence is known. In addition, there exist sequences that can easily be amplified and sequences that can hardly be amplified in the process of PCR reaction, and thus, the
35 efficiency of reaction (rate of amplification) is not uniform. For example, in the case where the content of a specific mRNA

5 in the total amount of extracted mRNA is examined to determine diseases or like based on the content, standard samples providing criteria should be always prepared to make correction on the above -described amplification rate.

10 Although the amount of the specimen solution required for a hybridization reaction decreases as the size of the substrate is reduced, there is a limitation on downsizing of the substrate in association with handling. Specifically, it is possible in principle to enhance array density and reduce the number of probes to be placed on the array to downsize the
15 substrate, but if an extremely small substrate is used, a dedicated handling apparatus is required in the process of processing such as hybridization reaction and detection thereafter, which cannot be practical.

Also, for examining cDNA for mRNA that is transcribed
20 with reflection of the process of development of a certain organism, cDNA for mRNA that is transcribed with reflection of each phase in the process of culturing a certain cell, cDNA for mRNA that is transcribed by interaction with drugs, and so on, a DNA array with multiple types of test samples arranged
25 is used. Examples of arraying this test sample are described, for example, in the above -described "Science", Vol. 270, pp 467, (1995). In this case, test samples arrayed on the substrate are dipped using as a probe solution the labeled DNA of known sequence that is derived from genes having a specific
30 function, whereby a hybridization reaction is carried out.

If a plurality of items is to be examined at a time using this methodology, DNA probes labeled with different types of fluorescent reagents (fluorochromes) should be prepared depending on the number of items. When a detection is made,
35 those different types of fluorescent reagents (fluorochromes) must be ~~detected as~~ distinguished from one another, and

5 therefore, their wavelengths and the like should be different
as a matter of course. Of course, detection filters
corresponding to respective fluorescent reagents
(fluorochromes) are also needed for a detector.

This need for a simultaneous examination of multiple
10 items for multiple specimens is not characteristic exclusively
of a hybridization reaction among genes (DNA).

For example, it is also important to examine multiple
items with a small amount of samples as to the interaction
between genes and other substances, such as interaction
15 between genes and proteins (DNA binding proteins) and
screening of chemicals that are bound to genes. Detection of
former DNA binding proteins is used to elucidate the control
mechanism of gene expression by proteins, such as
transcription accelerators ~~but~~. However, in the present
20 situation, methods in which DNA fragments are bound to
proteins, and thereafter complexes are analyzed by gel
electrophoresis, are adopted. In this method, the number of
specimens that can be analyzed at a time is limited due to the
usage of gel electrophoresis, and considerable time is
25 required for analysis.

For the field of drug development ~~of drugs~~, there may be
cases where an examination of interaction between genes and
administered drugs constitutes an important item in the
progress of research, but it takes relatively ~~much~~ a large
30 amount of time and ~~efforts~~ effort to obtain chemically
synthesized products for use in drugs to be researched, and it
can be considered that a reduction in the amount of drugs to
be used in screening results ~~in~~ is a significant improvement in
~~efficient~~ efficiency of their research.

35 As introduced above, there are cases where when a complex
is formed using an interaction between two substances, such as

5 hybridization between DNAs, formation of a complex of DNA and
a protein, and interaction of a drug compound with gene DNA,
or the presence or absence of the interaction causing a
complex to be formed, is examined, the amount of samples of
one of those two substances is limited, and the limited amount
10 of samples should be used to conduct a series of desired
examinations across multiple types as to the presence or
absence of formed complexes. That is, development of an
examination method in which consumption of samples required
for individual examinations can be reduced to carry out the
15 examination across multiple types more efficiently within a
limited amount of samples is desired.

SUMMARY OF THE INVENTION

An object of the first invention is to provide a method
of examining multiple specimens at a time for multiple items,
20 for example a method in which matrix substrates with
biological samples having different properties and origins
bound thereto are prepared, and on each matrix region,
oligonucleotides or proteins having different sequences and
drugs are spotted in an array form, whereby multiple specimens
25 are examined at a time for multiple items.

Another object of the invention is to provide a method in
which multiple specimens can also be examined at a time for
multiple items in a similar way for interaction between
chemicals, especially drugs, and cDNA, binding of proteins to
30 cDNA and the like.

An object of the second invention is to provide a new
method in which an oligonucleotide ~~of which~~ with a known base
sequence ~~is known~~ and which can be obtained relatively easily
is used as a detection probe ~~and when~~. When, for a limited
35 amount of sampled specimens, the presence or absence of a

5 bonding capability to the above _described oligonucleotide as
a detection probe or the degree of the bonding capability is
examined by the presence or absence of complexes formed
between those two substances, or efficiency thereof is
evaluated, consumption of specimens required for evaluation
10 for each type of oligonucleotide as a detection probe can be
reduced. In addition, the invention also has an object to
provide a detection substrate with the above _described
oligonucleotide being fixed as a detection probe in a
predetermined region of its surface, which is used exclusively
15 for the method, and to provide a method of preparing the
detection substrate.

The examination method of the first invention capable of
achieving the above _described objects is a method in which a
reactivity between a first sample and a plurality of second
20 samples having different properties from one another is
examined at a time,

characterized in that in a defined region on a substrate
with the first sample bound on the entire surface in advance,
the second samples are placed independently of one another as
25 spots having a smaller size than the above _described defined
region, and then the reactivity between the above _described
first sample and each of the second samples is tested.

The matrix of biological samples related to the invention
that is usefully used for the above examination method is
30 characterized in that two or more types of biological samples
of different origins exist in respective matrix regions
separated on the substrate.

According to the invention, a substrate with biological
samples having different properties and origins (e.g. nucleic
35 acids and proteins) bound in a matrix form in advance can be
provided.

5 There is also provided a method in which DNA probes like
oligonucleotides, cDNAs, proteins or chemicals are spotted in
an array form on the above -described substrate with
biological samples having different properties and origins
placed in a matrix form to carry out the reaction, and the
10 presence or absence of another sample bound to a certain
biological sample, the degree of the bonding, and the presence
or absence of interaction is quickly examined for multiple
items at ~~a the same time and speedily~~.

 In this method, the area occupied by one specimen is very
15 small, because two or more types of specimens are placed on
one substrate. Therefore, there is an advantage in that the
amount of required cDNA may be very small as compared to the
case where the hybridization reaction is carried out using a
conventional DNA array with an enormously large number of DNA
20 probes bound in an array form in advance. Also, there is
neither a limitation on the size of the DNA array substrate
nor an inconvenience ~~for~~ in handling.

 Also, by providing a method in which examination can be
carried out even with a small amount of samples, the method
25 opens the door to areas in which examination could not be
carried out, because conventionally, a sufficient amount of
samples cannot be obtained, for example, a new examination
area in which mRNA obtained from tissues is directly examined.

 In addition, according to the invention, ~~a method in~~
30 ~~which~~ chemicals, proteins and nucleic acids can be examined at
a the same time under the same reaction condition on the same
substrate.

 A method of detecting object components in test samples
according to the second aspect of the invention is a method in
35 which using as a detection probe oligonucleotide ~~of which~~ with
a known base sequence ~~is known~~, complexes formed between the

5 above _described oligonucleotide and the object components are detected to examine whether or not the object components ~~having a capability~~capable of binding to the above _described oligonucleotide are contained in the liquid test samples, or evaluate the degree of binding capability thereof,

10 characterized in that there is at least one type of the above _described oligonucleotide used as a detection probe, ~~of which with a known base sequence is known,~~

there are at least two types of test samples to be examined, and

15 a detection substrate with the above _described one or more types of oligonucleotide for detection probes bound to predetermined sections respectively on a predetermined solid substrate is used~~7.~~

~~the~~The above _described method ~~comprising~~comprises steps
20 of:

spotting a plurality of predetermined amounts of sample solution for each spot so that a predetermined array shape is formed in the spotted position, for each of the above _described two or more types of test samples, in each section
25 with the oligonucleotide ~~for detection~~to detect probes bound in advance;

detecting the presence or absence of complexes formed between the above _described oligonucleotide and the object component, for the above _described plurality of spots for
30 each test sample, respectively; and

determining whether or not the object component ~~having a capability~~capable of binding to the above _described oligonucleotide is contained, or the degree of the capability of binding, based on the result of the above _described
35 detection.

5 Also, the present invention provides a detection
substrate that is exclusively used when the above _described
method of the invention is carried out. That is, the
detection substrate of the present invention is a detection
substrate with two or more oligonucleotides having known base
10 sequences different from one another fixed on a solid
substrate, characterized in that:

the above _described plurality of oligonucleotides are
bound and fixed in predetermined sections, respectively, so
that one type of oligonucleotide exists in each section, and
15 a plurality of the above _described sections in which
oligonucleotides are fixed is placed in a matrix form on the
surface of the above _described solid substrate.

The method of preparing the detection substrate of the
present invention is a method suitable for preparation of the
20 above _described detection substrate of the invention, and
specifically, _ is a method of preparing a detection substrate
with two or more oligonucleotides having known base sequences
different from one another fixed on a solid substrate,
characterized in that:

25 for the above _described solid substrate, a substrate
with a plurality of sections separated in a matrix form in
advance formed on the surface thereof is used,

the above _described ~~a~~ plurality of oligonucleotides is
supplied into predetermined sections in predetermined amounts
30 using printing by an ink jet process, respectively, so that
one oligonucleotide is present in each section, and

the supplied oligonucleotides are fixed in the
predetermined sections.

5 BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 shows one example of an arrangement aspect of defined regions on a substrate in the present invention;

FIGS. 2A and 2B show one example of matrices in the present invention, wherein FIG. 2A is a plan view, and FIG. 2B
10 is a 2B-2B sectional view thereof;

FIG. 3 is a schematic explanatory view of a specimen solution discharging method by a bubble jet process that is an embodiment of the present invention;

FIG. 4 is a sectional view of a bubble jet head 105 taken
15 in the 4-4 line in FIG. 3;

FIG. 5 shows a layout of 64 discharged DNA probes on each black matrix;

FIG. 6 shows one example of detection substrates of the present invention, illustrating schematically a situation in
20 which sections in which oligonucleotides being detection probes are fixed are arranged in a matrix form, and a plurality of cDNAs are spotted in a two-dimensional array form onto each section as detection samples;

FIG. 7 illustrates schematically arrangements of
25 respective probes in the detection substrate with 64 DNA probes bound to sections arranged in the form of a 8 x 8 matrix, respectively;

FIG. 8 shows schematically a pattern of a spot ~~array of total-64 x 64~~ array in which 64 test samples are spotted in
30 the form of a two-dimensional 8 x 8 array on each section, for the detection substrate on which sections with probes fixed therein are arranged in the form of the 8 x 8 matrix;

FIG. 9 shows schematically a result of spotting 64 test samples in the form of the two-dimensional 8 x 8 array on each
35 section for 64 probes fixed in sections arranged in the form

5 of the 8 x 8 matrix to carry out the hybridization reaction;
and

FIG. 10 shows an example of the structure of sections
delimited by hydrophobic frame-structured walls provided on
the detection substrate of the present invention, and arranged
10 in the form of the 8 x 8 matrix.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

One embodiment of the present invention will be described
below referring to FIG. 1. FIG. 1 shows a substrate surface
with 64 defined regions formed thereon, wherein each region
15 (matrix) measures 1 mm by 1 mm, and a space x between regions
can be selected freely. For methods of preparing biological
sample binding matrix substrates, for example, a method can be
used in which the solution of a first sample (e.g. biological
sample) is printed on the entire surface of defined regions on
20 the substrate as a "solid print pattern" by coating and ink
jet processes, or is supplied by methods, such as chemical
synthesis on the substrate, and is bound in a matrix form on
the substrate through adsorption to the substrate or chemical
reaction between functional groups existing in the biological
25 sample and functional groups existing on the substrate.
Furthermore, the situation in which the first sample is bound
on the entire surface of defined regions means a situation in
which the first sample is bound across the entire surface,
such that when a second sample and samples thereafter are
30 supplied in these defined regions, these reactions occur
without being limited to the positions in the above _described
regions in which the samples are supplied. For example, the
first sample may be fixed in layered form on the entire
surface, or the masses of molecules constituting the first

5 sample may be dispersed on the entire surface in high density with micro-spaces being kept among them.

The defined regions on this substrate may previously be provided on the substrate as a well constituted by sections separated in a pattern ~~form~~formed by walls of hydrophobic
10 compounds.

Also, when using a substrate with ~~nucleosidea~~ nucleic acid (cDNA) being a biological sample fixed thereon as the first sample, two or more probe DNAs possibly included in cDNA are contacted with cDNA on the substrate as the second sample
15 and samples thereafter, ~~and products.~~ Products of the reaction with the above -described probes are detected on the above -described solid phase to detect the presence or absence of probe DNA sequences in the above -described cDNA, two or more probes are supplied in an array form as mutually
20 independent spots in each matrix with various kinds of cDNA bound in the defined regions, thereby making it possible to perform simultaneous detection with two or more probes.

Also, on the nucleic acid (cDNA) matrix, two or more types of chemicals or proteins that are possibly bound to cDNA
25 are contacted with the probe DNA on the substrate as mutually independent spots, thereby making it possible to perform a multiple-item examination composed of these reactions at ~~a~~the same time. Multiple-item screening of DNA binding proteins and DNA binding chemicals can be performed at ~~a~~the same time
30 by detecting the presence of binding of chemicals or proteins to probes on the solid phase.

The present invention is characterized by supplying probe DNA, proteins and chemicals in a form of droplets of small amounts on the matrix on which biological samples, such as
35 cDNA are applied, wherein different types of samples are

5 arranged in an array form, thereby making it possible to perform simultaneous multiple-item processing.

Combinations of the first sample fixed in advance on the substrate and the second sample and sample thereafter that are reacted with the first sample may include the following
10 combinations.

Specific examples of the matrix or the like formed of defined regions on the substrate for use in the present invention will be described below.

~~(Shapes of matrices with biological samples bound thereto)~~
15 Matrices with Biological Samples Bound Thereto

The shapes of matrix patterns are not particularly limited, ~~and may include any shapes, but shapes such as~~ but linear, ~~squares~~ square and rectangular shapes are ~~preferable~~ preferred in that they can be treated irrespective
20 of how specimens are supplied, ~~in consideration of convenience at the time of supplying specimens on the created substrate.~~ Of course, ~~forms~~ shapes such as circles and ellipses will cause no problems.

Materials that are fixed to the substrate as a first
25 sample may include unknown base sequences derived from organisms, cDNA libraries, mRNA libraries, sets of two or more DNA and RNA, known DNA and RNA synthesized or derived from organisms or sets thereof, chips of cloned oncogenes, protein fractions including at least one type of protein derived from
30 organisms, proteins of single type, mixtures of known proteins of different types, and chemicals.

~~(Density of matrices with biological samples bound)~~
Matrices with Bound Biological Samples

The density of matrices is not particularly limited, but
35 for a preferred form, the density of 400 per centimeter square is preferable. For this preferred density of 400/cm², the

5 size of one matrix is a 500 μm square in the case of a square
~~formshape~~. If samples to be arranged as spots on the array
are ~~arranged as spots with diameters of 100 μm in diameter~~, 25
spots are arranged in total ~~with~~, 5 spots high by 5 spots
wide. Also, if the diameter of sample solution is 20 μm , the
10 number of spots that can be arranged in a row is 25, and 625
spots can be arranged in total.

~~(Preparation of a substrate with biological samples bound~~
~~thereto)~~ Substrate with Biological Samples Bound Thereto

Samples originated from organisms (biological samples)
15 include nucleic acids and proteins. Nucleic~~7~~ acids include,
for example, mRNA and cDNA, and methods for binding them on
the substrate include a method in which a nucleic acid
extracted and purified in advance is applied to the substrate
to fix the nucleic acid by adsorption and an electrostatic
20 bond, and a method in which the nucleic acid is fixed by
providing a covalent bond ~~through~~ through a chemical reaction
with functional groups on the substrate using amino groups of
the nucleic acid ~~has~~.

The method using negative electric charges of DNA is a
25 method in which the nucleic acid is electrostatically bound to
a solid carrier subjected to a surface treatment with ~~poly~~
positive polymeric ions, such as polylysine, polyethyleneimine
and polyalkylamine, and then blocking of excessive positive
ions is carried out, which is generally used.

30 ~~(Types of functional groups of solid phases and nucleic~~
~~acids)~~ Types of Functional Groups of Solid Phases and Nucleic
Acids

Combinations of functional groups that are used for
fixation include, for example, a combination of epoxy groups
35 (on solid phase) and amino groups (amino groups in nucleic
acid probe terminals or base groups). Methods for introducing

5 epoxy groups to the solid surface include, for example, a
method in which polyglycidyl methacrylate having epoxy groups
is applied to the solid surface composed of resin, and a
method in which a silane coupling agent having epoxy groups is
applied to the solid surface made of glass and is reacted with
10 glass.

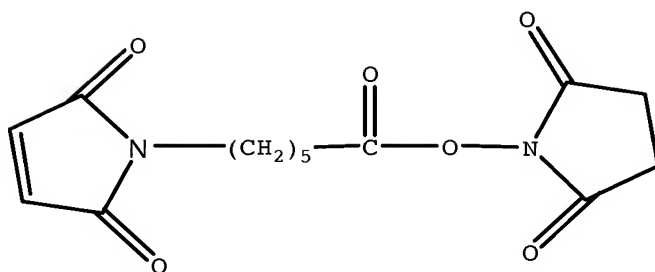
~~(Binding of ~~proteins~~Proteins to the ~~solid phase~~)Solid~~
Phase

Methods of binding proteins to the substrate include
methods using adsorption, as in the case of nucleic acid, and
15 methods using electrostatic binding. Furthermore, methods of
forming a covalent bond include ~~methods~~those using SH groups
of cysteine residues in addition to the above described
methods using amino groups.

~~(Methods of ~~fixation of proteins using thiol~~~~
20 ~~groups)~~Fixing Proteins Using Thiol Groups

Methods using cysteine residues for ~~fixation of~~fixing
proteins include, for example, methods using combinations of
maleimide groups and thiol groups (-SH). That is, the
treatment is ~~done~~performed so that the solid surface has
25 maleimide groups, whereby thiol groups of cysteine residues
supplied to the solid surface can be reacted with maleimide
groups of the solid surface to fix proteins.

~~For methods of introducing maleimide groups to the solid~~
~~surface, a various~~Various kinds of methods may be used, ~~and~~
30 ~~this~~to introduce maleimide groups to the solid surface. This
can be achieved by, for example, reacting an aminosilane
coupling agent with a glass substrate, and then reacting its
amino groups with a reagent containing N-(6-
maleimidocaproyloxy) succinimide) expressed by the following
35 structural formula (EMCS reagent: manufactured by Dojin Co.,
Ltd.)



5

~~For~~As another example, a reagent containing succinimidyl 4-(maleimidophenyl) butyrate can be used to react with ~~amino groups~~, preferably amino groups.

~~(DNA matrix structures composed of hydrophobic matrices)~~

10

DNA Matrix Structures Composed of Hydrophobic Matrices

~~For~~As an additional form of ~~fixation of~~fixing biological samples, a method can be used in which a well composed of, for example, hydrophilic and hydrophobic matrices is formed on the solid surface, a structure to prevent coupling among spots is provided in advance, and the DNA prove is supplied in the well to carry out a coupling reaction.

15

~~(Materials of matrices/wells)~~Matrices/Wells

When a prove solution is put on the separated matrix to carry out the coupling reaction, it is preferable that portions constituting the well ~~is~~are hydrophilic, and portions corresponding to the wall surface of the well and the partition between the well and a neighboring well are composed of materials whose surfaces are less compatible with the prove solution. Due to such a treatment, the probe solution can be smoothly supplied to a desired well even if some positional deviation occurs when the prove solution is supplied to the well.

20

25

One example of ~~matrices~~a matrix in this embodiment is shown in FIGS. 2A and 2B. FIG. 2A is a plan view, and FIG. 2B is a 2B-2B sectional view thereof. This matrix has a structure in which a matrix ~~patter~~pattern 125 having a frame structure with formed recesses 127 (wells) placed in the form of a solid phase 103 is provided. The wells 127 separated from one another by the matrix 125 (height) are provided as

30

5 through-holes (cut-off portions) in the matrix pattern, ~~of~~
~~which.~~ The side of the matrix pattern is constituted by
~~heights~~height, and ~~of which the~~ bottom 129 has the exposed
surface of the solid phase 103. The portion of the exposed
10 surface of the solid phase 103 forms a surface that can be
coupled to the probe, and the probe is fixed in a
predetermined recess.

Materials forming the matrix pattern include, for
example, metals (chrome, aluminum, gold, etc.) and resins.
They include resins, such as acryl, polycarbonate,
15 polystyrene, polyimide, acrylate monomers and urethane
acrylate, and photosensitive resins, such as photoresists
having black dies and black pigments contained therein. For
specific examples of photosensitive resins, UV resists, DEEP-
UV resists, ultraviolet cured resins and the like can be used.
20 UV resists may include negative resists, such as cyclized
polyisoprene-aromatic ~~pi~~isazide~~bisazide~~ resists, phenol resin-
aromatic azide compound resists, and positive resists, such as
novolac resin-diazonaphthoquinone resists.

DEEP-UV resists may include, for example, radiation
25 dispersion type polymer resists, such as polymethyl
methacrylate, polymethylene sulfone, polyhexafluorobutyl
methacrylate, polymethyl ~~isopropenyl~~isopropenyl ketone and
bromo poly 1- trimethylcylilpropine, and dissolution
inhibiting resists, such as cholate o-nitrobenzyl ester as
30 positive type resists, and may include
~~polyvinylpheno~~polyvinylphenol-3-3'-diazidediphenylsulfone,
and polymethacrylate glycidyl as negative type resists.

Ultraviolet cured resins may include polyester acrylate,
epoxy acrylate and urethane diacrylate containing
35 approximately 2 to 10% by weight of one or more types of
photopolymerization initiators, which are selected from
benzophenone and substituted derivatives thereof, oxime
compounds, such as benzyl, and so on.

For curbing reflection by the material forming the matrix
40 during detection, light-blocking materials can be effectively

5 used ~~for materials forming~~to form the matrix pattern. For this purpose, it is effective to add black pigments in the above described resin, and for. Examples of black pigments, that can be used are carbon black and black organic pigments ~~can be used.~~

10 Here, if the matrix 125 is composed of resin, the surface of the matrix 125 is hydrophobic. This structure is preferred when an aqueous solution is used as a solution containing probes to be supplied to the well. That is, even if the probe solution is supplied to the well, the probe solution is
15 supplied to a desired well quite smoothly. Also, if different probes are supplied among adjacent wells at the same time, intermingling (cross contamination) of different probe solutions supplied among these wells can be prevented.

The thickness of the matrix (height from the solid
20 surface) is determined in ~~the~~ light of the matrix pattern forming process and the volume of the well, but it is preferably in the range of 1 to 20 μm . Particularly, it can be considered as a thickness range allowing that effectively prevents cross contamination ~~to be prevented effectively~~ when
25 the probe solution is supplied to each well ~~though~~through an ink jet process.

~~(Types of samples~~Samples to be spotted)Spotted

Samples to be spotted as droplets onto the above described matrices of biological samples include probe nucleic acids, proteins and chemicals, such as drugs.
30

For probe nucleic acids, in addition to deoxyribonucleic acid, any ~~type~~type of nucleic ~~acids~~acid, such as ribonucleic acid and peptide nucleic acid, may be used as long as ~~they~~ have it has nucleic acid bases. The length of the
35 oligonucleotide probe is not particularly limited, but it is preferably in the range of 10 mer to 50 mer for carrying out an accurate hybridization reaction with cDNA.

For proteins, their own fluorescence can be used to detect DNA bonding proteins.

5 Some chemicals can also be detected with their own
fluorescence.

~~(Method of preparing sample arrays)~~Preparing Sample
Arrays

10 Methods of spotting sample solution on defined positions
in the size of several tens to several hundreds of microns
include a pin system, an ink jet system and a capillary
system.

15 The pin system refers to a method in which the sample is
attached to the pin tip, for example, in such a manner that
the pin tip is contacted with the surface of the solution
including the sample, and then the tip is mechanically
contacted with the solid phase, thereby preparing a sample
array. The capillary system using a capillary is such that
the sample solution once sucked up to the capillary is
20 mechanically contacted with the solid phase through the tip of
the capillary as in the case of the pin system, thereby
supplying the sample solution in an array form. For these
spotting operations, various ~~apparatuses~~ commercially
available apparatuses from various companies may be used.
25 These methods are considered as most preferable ~~methods~~ in the
sense that any sample DNA can be supplied. However, as for
quantification, the problem may be ~~unsolved~~ in that viscosity
varies depending on the length and concentration of DNA. For
proteins, these methods are also preferred in the sense that
30 they are deposited independently of the size and viscosity of
molecules, but not suitable for ~~quantitative~~ quantitative
analysis.

~~(Outline of sample array preparing methods through the
ink jet process)~~

35 Outline of Sample Array Preparing Methods Through the Ink
Jet Process

 Samples capable of being discharged in an ink jet process
include chemicals in addition to nucleic acids and proteins.

40 In the ink jet process, because a shearing force is
exerted, the length of dischargeable nucleic acids and the

5 size of dischargeable proteins are often limited. However,
~~it~~this process is superior in quantification to the pin system
and capillary system, and is used more suitably than other
systems with respect to the discharge of chemicals.

Dischargeable nucleic acids are limited to those with a
10 relative length to bases of 5 kb or smaller, and dischargeable
proteins are limited to those of 1000 K daltons or less. As
for chemicals, any chemicals can be discharged.

~~For liquids for discharge to~~Any liquid can be used for
discharging and supplying samples with ink jets, ~~any liquid~~
15 ~~can be used~~ as long as ~~it~~this liquid is capable of being
discharged from ink jets, ~~and the.~~ The above -described
liquid discharged from the head is shot into a predetermined
position, ~~and in the state of.~~ When being mixed with nucleic
acid probes and during discharge, the above -described nucleic
20 acid probes are not damaged.

~~And, in~~In terms of dischargeability from the ink jet,
particularly from the bubble jet head, ~~for~~with respect to the
properties of the above -described liquid, it is preferable
that its viscosity isbe in the range of 1 to 15 cps and its
25 surface tension isbe 30 dyn/cm or larger. Also, if the
viscosity is in the range of 1 to 5 cps and the surface
tension is in the range of 30 to 50 dyn/cm, the
~~position~~positions in which the liquid is spotted on the solid
phase isare extremely accurate, ~~allowing the method to be used~~
30 ~~particularly suitably.~~

Therefore, if the stability of the nucleic acid during
discharge or the like is taken into consideration, a nucleic
acid probe of, for example, 2 to 5000 mer, particularly 2 to
1000 mer, is preferably contained in the solution ~~in~~
35 ~~concentrations~~at a concentration of 0.05 to 500 μ M,
particularly 2 to 50 μ M.

FIG. 3 is a schematic explanatory view of a specimen
solution discharging method through the bubble jet process
~~that,~~ which is one embodiment of the present invention. In
40 FIG. 3, reference numeral 101 denotes a liquid supplying

5 system (nozzle) retaining a solution, including a specimen, as
a discharge liquid in such a manner that the solution is
capable of being discharged, reference numeral 103 denotes a
solid phase having a nucleic probe bound thereto with which
the above -described specimen is reacted, and reference
10 numeral 105 denotes a bubble jet head ~~having a function of~~
~~giving that~~ supplies heat energy to the above -described liquid
to discharge it, ~~which and~~ is a type of ink jet head.
Reference numeral 104 denotes a liquid including the specimen
discharged from the bubble jet head. FIG. 4 is a 4-4 line
15 sectional view of the bubble jet head 105 in FIG. 3, ~~and in 3.~~
In FIG. 4, reference numeral 105 denotes the bubble jet head,
~~and reference.~~ Reference numeral 107 denotes a liquid
including a specimen solution to be discharged, ~~and reference.~~
Reference numeral 117 denotes a substrate portion having a
20 heat generation portion to ~~give~~ provide discharge energy to the
above -described liquid. The substrate portion 117 includes a
protective layer 109 formed ~~by from~~ silicon oxide and the like,
electrodes 111-1 and 111-2 formed ~~by from~~ aluminum and the
like, an exothermic resistor layer 113 formed ~~by from~~ nichrome
25 and the like, a heat storage layer 115, and a support 116
formed ~~by from~~ aluminum having a good heat-release property.

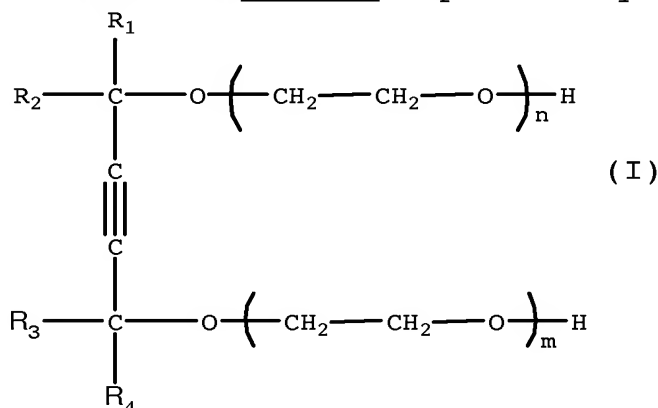
The liquid ~~107~~ 107, including the specimen, comes to a
discharge orifice (discharge outlet) 119, and forms a meniscus
121 with a predetermined pressure. Here, when electric
30 signals are applied to the electrodes 111-1 and 111-2, a
region (foaming region) denoted by reference numeral 123
abruptly releases heat, and the liquid 117 contacted therewith
is discharged and flies toward the solid surface 103. The
amount of the liquid that can be discharged using a bubble jet
35 head having such a structure varies depending on the size of
its nozzle, but can be controlled approximately to 4 to 50
picoliters, which is extremely useful as a means for placing
specimen probes ~~in at~~ a high density.

~~And, in~~ In terms of dischargeability from the ink jet,
40 particularly from the bubble jet head, ~~for~~ with respect to the

5 properties of the above described liquid, it is preferable
that its viscosity ~~is be~~ in the range of 1 to 15 cps and its
surface tension ~~is be~~ 30 dyn/cm or larger. Also, if the
viscosity is in the range of 1 to 5 cps and the surface
tension is in the range of 30 to 50 dyn/cm, the positions in
10 which the liquid is spotted in an exceedingly accurate
position on the solid phase, allowing the method to be used
particularly suitably are extremely accurate.

Therefore, if the stability of nucleic acid during
discharge or the like is taken into consideration, a nucleic
15 acid of, for example, 2 to 5000 mer, particularly 2 to 1000
mer, is preferably contained in the solution ~~in~~
concentrations at a concentration of 0.05 to 500 μ M,
particularly 2 to 50 μ M.

~~For the~~The composition of a discharged liquid, the
20 ~~composition of~~ liquid is not particularly limited, as long as
the liquid has no substantial influence on the nucleic acid
probe when it is mixed with the nucleic acid probe and when it
is discharged from the ink jet, and it can be normally
discharged to the solid phase using the ink jet, ~~but~~
25 ~~preferable are.~~ Preferable liquids including include glycerin,
urea, thiodiglycol or ethylene glycol, isopropyl alcohol, and
acetyl ~~alcohol~~ alcohol expressed by the following formula:



30 In the above formula (I), R_1 , R_2 , R_3 and R_4 represent
alkyl groups, specifically linear or branched alkyl groups
having 1 to 4 carbon atoms, m and n represent integer numbers,

5 respectively, wherein m and n equal 0, or $1 \leq m + n \leq 30$ holds, ~~30~~,
and if $m + n = 1$ holds, ~~1~~, m or n equals ~~0~~ 0.

~~Further~~More specifically, a liquid containing 5 to 10% by
weight (wt%) of urea, 5 to 10 wt% of glycerin, 5 to 10 wt% of
thiodiglycol, and 0.02 to 5 wt%, more preferably 0.5 to 1 wt%,
10 of acetylene alcohol presented by the above formula (I) ~~is~~may
be suitably used.

The detecting method of the present invention is a method
of detecting a complex formed between an oligonucleotide for
detection probes and an object component, which is used for
15 the purpose of making an evaluation/examination as to whether
or not a component ~~having capability~~capable of binding to the
oligonucleotide and forming a complex in a liquid test sample
for use as a detection probe whose base sequence is known, and
~~forming therewith a complex exists in a liquid test sample,~~
20 ~~and~~ as to the degree of binding capability thereof if such a
component exists in the sample. For detecting this complex,
the oligonucleotide for detection probes is fixed in advance
on the solid surface substrate, whereby this fixed
oligonucleotide is bound to the object component contained in
25 the test sample, and the formed complex is separated while it
is fixed on the solid substrate, ~~and on~~. On the basis of a
methodology for detecting complexes using proper detecting
means, the amount of test samples required at this time is
reduced to a very low level, ~~and also~~. Also, the detection
30 accuracy and ~~sensitiveness~~sensitivity are kept at a
sufficiently high level.

That is, in this methodology providing a base for the
present invention, since surface density of the
oligonucleotide for detection probes that is fixed on the
35 solid surface substrate can be kept at a predetermined value,
the amount of the formed complex is proportional to the

5 binding capability of the object component, and is also
proportional to the concentration of the object component
contained in the test sample that is contacted with the solid
surface substrate and is made to act on the oligonucleotide.
Taking advantage of this characteristic, the test sample is
10 contacted only with the surface, with the oligonucleotide for
detection probes actually fixed thereon, ~~and the.~~ The contact
area is limited to a certain ~~level~~degree, whereby the amount
of the used test sample is also limited to a certain
~~level~~degree. Specifically, by adopting a means for spotting a
15 predetermined minimal amount of liquid in the form of
droplets, the contact area and the amount of liquid put
thereon ~~is~~are controlled with good reproducibility. The
amount of the complex that would be automatically fixed on the
solid surface substrate with formation is detected for this
20 limited contact area, thereby achieving detection accuracy and
~~sensitiveness~~sensitivity that are essentially as high as those
in the case of dipping ~~in the liquid test sample the whole of~~
~~the~~the entire solid substrate with oligonucleotide for
detection probes fixed thereon in the liquid test sample.

25 The complex is detected by the label bound on the surface
of the substrate. When a complex of the oligonucleotide and
the labeled test sample is formed and the individual spots are
sufficiently spaced from each other, detection can be carried
out independently for each spot. Therefore, if given or
30 larger spaces are provided between adjacent spots, even though
there are spots for different test samples nearby, only spots
for desired test samples can be selected to continue detection
work without being influenced by those spots. In the
detection method of the present invention, in order to satisfy
35 reliably ~~this~~, the requirement that given or larger spaces be
provided between adjacent spots, a defined array is formed in

5 the spot position as a result of providing predetermined spaces as spaces between spots, and a predetermined amount of sample solution is spotted for each spot to make the spot area (contact area) constant, or make the spot diameter constant to ensure reproducibility, because the shape of the spot (contact
10 surface) is generally a circle. As a matter of course, for precluding the influence of adjacent spots, a space between spots is selected such that optical signals (fluorescent) and the like derived from the adjacent spots are not mixed in the detection system, in the light of the measured area (diameter
15 in the measurement range) of the detection system selected as appropriate in accordance with the spot diameter. Also, as a matter of course, the detecting method of the present invention really shows its advantages in the case where there exist two or more types of test samples, and they are detected
20 simultaneously.

On the other hand, on the surface to which a plurality of spots of such array forms ~~is~~are provided, one type of oligonucleotide for detection probes should be fixed ~~in~~at a uniform surface density. Also, for the section in which the
25 oligonucleotide for detection probes is fixed, its area and shape are selected as appropriate in accordance with the above -described array space and the total number of spots to be included in a series of arrays. It is also possible to provide sections having different oligonucleotides fixed
30 therein in different regions on the detection substrate to be used, and to place a plurality of sections with two or more oligonucleotides fixed therein, respectively. That is, it can be said that the detecting method of the present invention becomes a more suitable method if used when two or more types
35 of nucleotides are used as detection probes to carry out a series of evaluations simultaneously for a plurality of test

5 samples, with respect to two or more types of object components corresponding to respective oligonucleotides.

Generally, in such an evaluation, it is often the case ~~not~~ that the oligonucleotides for detection probes are predetermined while only an approximate number of test samples
10 to be evaluated is determined. In such a case, it is preferable that as a detection substrate with oligonucleotides for detection probes fixed thereon in advance, a detection substrate with two or more types of detection probes put thereon systematically, having on the substrate surface in a
15 matrix form sections in which respective oligonucleotides are fixed. In this detection substrate with fixed sections arranged thereon in a matrix form, the unit of a total number of spots that are made in an array form in each section is fixed, but a plurality of these units ~~of number of spots~~ can
20 be used to carry out the evaluation depending on the number of test samples to be actually evaluated, thus enhancing convenience in practice. Furthermore, for ~~the~~ each section arranged in a matrix form, a pattern formed by hydrophobic compounds is preferably provided in its ~~substrate~~ substrate
25 to provide a form in which mutual regions are separated from one another.

In the detecting method of the present invention, nucleic acid molecules may be selected as object components to apply the same ~~to evaluation as to~~ when evaluating whether or not
30 they are engaged in a double-strand formation ~~into~~ in hybrid substances formed through a hybridization reaction with the oligonucleotide for detection probes. In this case, the method used is an effective method in which an evaluation is made at ~~a~~ the same time even for multiple test samples, as to
35 whether or not nucleic acid molecules, including base sequences complementary to known base sequences ~~that~~ of the

5 oligonucleotide for detection probes ~~has~~, are contained in the
test sample. Alternatively, if two or more types of
nucleotides for detection probes are provided, and one type of
nucleic acid molecules ~~are~~is contained in each test sample, an
10 ~~evaluation can be made for the nucleic acid molecule of~~
~~which with the still unknown base sequences are still unknown,~~
as to whether or not the nucleic acid molecule includes base
sequences complementary to known base sequences ~~that of~~ each
oligonucleotide ~~has~~, which is effective, for example, ~~for~~as a
means for ~~making search~~searching for a gene group having a set
15 of homologies.

The detection substrate of the present invention is a DNA
probe substrate with oligonucleotides for use in probes
respectively bound to sections arranged in a matrix form in
advance, ~~and particularly.~~ Particularly for the substrate
20 ~~itself~~, the bottoms of sections separated by wells (walls) of
the frame structure matrix patterns formed in advance by
hydrophobic compounds are formed as hydrophilic surfaces,
thereby making the binding of oligonucleotide easier. Also,
by providing this hydrophobic wall, intermingling of DNA
25 probes among adjacent sections can be curbed more reliably.

Also, using these DNA probe substrates, the test sample
is spotted in an array form on the matrix of the
oligonucleotide to carry out the hybridization reaction,
thereby providing a means for checking quickly whether or not
30 nucleic acid molecules ~~having complementarity with~~
complementary sequences are included in each test sample for a
certain oligonucleotide probe.

In this method, since the number of test samples that are
used in the hybridization reaction is determined ~~depending~~
35 solely based on the number of spots, the size of the detection
substrate is not limited, ~~and by.~~ By using a substrate ~~of~~with

5 a large area, the section in which each probe is fixed can be
widened, and the necessity to ~~enhance~~increase density can be
eliminated. Thus, since the section in which each probe is
fixed can be ~~widened~~increased, a wide range of methods can be
used, such as methods in which a liquid containing probes is
10 applied to, or printed as, a "solid printed pattern" through
an ink jet process on defined regions on the substrate, or
methods in which chemical synthesis is carried on the
substrate, for means for binding of the probe/oligonucleotide
on the substrate.

15 Also, considering that the probe/oligonucleotide is less
expensive and easier to obtain than the test sample, no
significant problem arises even if the area of the region to
which oligonucleotide is bound is more or less increased, ~~and~~
~~in.~~ In this case, with respect to ~~a~~ various kinds of test
20 samples to be spotted, it is not necessary to always spot them
~~in at a~~ high density. Furthermore, when the test sample is
spotted in small amounts, the concentration of the object
component that is contained in the test sample is increased,
whereby the hybridization reaction can be accelerated, ~~thus~~
25 making it possible to perform ~~high~~a highly sensitive detection
for a short time. In addition, application of the detecting
method of the present invention will open the door to fields
that could not be considered previously, because a sufficient
amount of samples could not be obtained, for example a new
30 field in which mRNA obtained from tissues is directly
examined.

 Furthermore, information of reactivity in association
with the obtained hybridization reaction is analyzed/evaluated
in terms of existence/~~not-non~~-existence of complementarity to
35 various kinds of oligonucleotides/probes, with respect to
nucleic acid molecules contained in a specific test sample,

5 thereby making it possible to carry out a detection having that
also functions similar to ~~those of the~~ conventional DNA arrays
(hybridization reaction with multiple probes for one
specimen).

Furthermore, the detecting method of the present
10 invention provides a means for evaluating as object components
the interaction between chemicals, in particular drugs and
oligonucleotides, the bonding of proteins to oligonucleotides
and the like, ~~and therefore.~~ Therefore, it can also be used
as a means making it possible to examine for examining object
15 components included in the test sample for multiple items,
with respect to a large number of test samples. In addition,
it provides a means making it possible to carry out for
carrying out an examination on the same substrate at ~~at the same~~
time and under the same conditions even for object components
20 of different properties, such as chemicals, proteins and
nucleic acids.

The detecting method of the present invention and the
detection substrate for use exclusively therein will be
described further in detail below.

25 ~~In FIG. 6 is shown~~ shows an example of applying the
detecting method of the present invention to an embodiment in
which ~~using cDNA~~ is used as an object component, ~~a.~~ A hybrid
substance is formed through a hybridization reaction with an
oligonucleotide ~~of~~ with known base sequence that is used for
30 detection probes. In the detection substrate shown in FIG. 6,
a plurality of rectangular sections separated systematically
in a matrix form in advance ~~is~~ are provided on the solid
surface substrate ~~of~~ in a rectangular form. The rectangular
sections are each spatially isolated by matrix compartments
35 that are surrounding walls. DNA probes that are used for

5 hybridization/probes are each bound uniformly to the bottom surface of the rectangular sections.

Also, attached is an enlarged view showing schematically a situation in which a plurality of test samples including cDNA as an object component, for example two or more types of cDNA solutions prepared based on m-RNAs collected respectively
10 are spotted in the form of the two-dimensional array like a square matrix, in a section with the DNA probe fixed therein. The detection substrate, the detection probe, components to be detected and the like that are used in the detecting method of the present invention will be described further in detail.
15

~~(Oligonucleotides that are used)~~Used for detection probes)Detection Probes

In the detecting method of the present invention, a deoxyribonucleic acid can be used for an oligonucleotide that
20 is used for detection probes. In addition thereto, a ribonucleic acid, a peptide nucleic acid and the like can be used. Types thereof are not limited as long as they have desired base sequences, ~~and~~ are capable of being bound to other molecules in those portions, and ~~also as long as they~~
25 can be fixed on a solid substrate. Also, for portions excluding nucleic acid chains, those modified with non-nucleic acid atom groups and those having additional structures and ~~so on~~the like can also be used as long as the above -described requirements are satisfied.

30 Furthermore, for this oligonucleotide that is used for detection probes, a desired amount thereof should be artificially prepared or collected, and its base sequence itself should be known. However, its nucleic acid part should have at least two bases. Its base length is not limited in principle, but if the length exceeds that of 100 bases,
35 ~~difficulty~~it becomes significantly more significantdifficult

5 to use as its base length increases when ~~fixation~~fixing on the solid substrate is to be carried out. Therefore, ~~and~~ ~~therefore~~ the base length is preferably restricted to that of 100 bases or less.

For example, when this oligonucleotide is subjected to a
10 hybridization reaction with, for example, nucleic acid molecules ~~with the length of that~~ are more than 100 bases long, the length of the oligonucleotide is preferably at least 10 mer for obtaining sufficient bonding. On the other hand, if the length exceeds 50 mer, it is difficult to set conditions
15 for controlling the detection of mismatching, thus making it difficult to select and detect only those that are fully matched. Thus, in order to detect mutations, the length is preferably 60 mer or ~~smaller~~less.

Furthermore, the range of 10 mer to 60 mer is a
20 preferable range even when the oligonucleotide having desired base sequences, for example DNA is prepared through chemical synthesis.

~~(Shapes of sections with oligonucleotide fixed therein, which is arranged in a matrix form)~~

25 Shapes of Sections with Oligonucleotide Fixed Therein and Arranged in a Matrix Form

The shape of a section itself in which the oligonucleotide for detection probes is bound and fixed is not particularly limited. However, if ~~considering that a test~~
30 ~~sample~~sample is spotted in an array form on this section, generally a simpler shape rather than a complicated outside shape is preferably selected. In addition, ~~also~~ when the oligonucleotide is bound and fixed, generally, a simpler shape is preferably selected for providing a uniform surface density
35 in such a section, in terms of working efficiency and convenience. Specifically, rectangular forms, for example, line forms, squares and rectangles, are preferably adopted.

5 Of course, in principle, forms whose perimeters are formed by curves, such as circles and ellipses, do not cause any problems.

On the other hand, in the detection substrate of the present invention, when two or more oligonucleotides that are
10 used for detection probes are put on one substrate, sections in which they are fixed are preferably arranged in a matrix form, in ~~terms~~view of working efficiency and convenience. Also, preferably, the form and area of each section is unified, ~~and its area is also unified.~~

15 ~~(Density of sections arranged in a matrix form)~~ Sections Arranged in a Matrix Form

The density of sections arranged in a matrix form is selected as appropriate depending on the number of oligonucleotides that are put on the detection substrate at a
20 time. Moreover, ~~but~~ the density of 400 per centimeter square or less is preferable. If the density is 400/cm², and the form of each section is a square, the size of each section is a 500 μm square. If test samples are closely arranged in an array form as spots with diameters of 100 μm, 25 spots are
25 arranged in total with 5 spots high by 5 spots wide. Also, if the diameter of the spot is 20 μm, the number of spots that can be arranged in a row is 25, leading to 625 spots in total. Since the detecting method of the present invention has more significant advantages when there ~~are~~is a large number of test
30 samples and they are examined at ~~a~~the same time, the final object of the invention will be more satisfactorily achieved if the density of the section that is arranged is selected so that at least the upper limit of the number of test samples that can be spotted approximately equals the above -described
35 value.

For example, when the detecting method of the present invention is applied to test samples including cDNA, the number of test samples to be examined, specifically the total number of types of cDNA often is approximately as many as
40 3600. In this case, if the diameter of the spot is 100 μm,

5 the size of one section approximately equals a 6 mm square
when 60 spots are arranged in rows and columns, respectively.
Also, even if the diameter of the spot is 20 μm , the size of
one section should be a 1.2 mm square. In this way, in the
detection substrate for use in the detecting method of the
10 present invention, there are not a few cases of application
objects where the density of sections that are arranged in a
matrix form is preferably selected as 400 per centimeter
square or less.

Furthermore, in the detecting method of the present
15 invention, the test sample is spotted as droplets, and in the
case where the diameter of the spot is 100 μm , for example,
the amount of liquid required for the droplet of one spot is
about 25 picoliters. Even if the number of probes for use in
examination is selected as 400 (for example, the number of
20 sections of the matrix to be provided on the substrate is 400)
for this spot size, the total amount of liquid required for
the whole spots may be no more than 10 nanoliters for each
test sample, thus making it possible to carry out an objective
examination ~~items~~ with a minimal amount of liquid.

25 Also, in the conventional method, in which the detection
substrate is dipped in the solution of the test samples, the
amount of the required liquid ~~is dependent~~ depends on the size
of the substrate. Thus, and thus if the amount of the test
sample is essentially very small, the size of the substrate
30 should be reduced in accordance with the amount of liquid, and
it is essential to highly integrate probes that are fixed on
the substrate. On the other hand, in the detecting method of
the present invention, the size of the substrate itself can be
freely selected without allowing for the liquid amount of the
35 test sample. In addition, when the oligonucleotide that is
used for detection probes is fixed, the surface density should
be ~~uniformed~~ uniform as a matter of course. However, but it is
not necessary to highly integrate a plurality of probes to fix
them, ~~thus~~ making the fixing operation easier.

5 ~~(Fixation~~Fixing of oligonucleotideOligonucleotide on the
substrate)Substrate

As means for fixing the oligonucleotide that is used for
detection probes on the surface of the substrate, a method in
which the oligonucleotide separately prepared in advance is
10 supplied in predetermined sections by coating or printing to
bind the oligonucleotide, or a method in which each
oligonucleotide, specifically, a DNA probe or the like is
synthesized in a solid phase on the substrate to prepare
originally bound DNA, can be used. Furthermore, even in the
15 case where the oligonucleotide is not DNA, but, for example,
is ribonucleic acid or peptide nucleic acid, synthesis on the
substrate can be carried out to bind the oligonucleotide as
described ~~later~~below.

On the other hand, when the oligonucleotide, specifically
20 DNA or ribonucleic acid, peptide nucleic acid or the like,
separately synthesized or collected in advance, is used for
detection probes, a process of fixing the oligonucleotide by
covalent ~~bond~~bonding or ~~of fixing it by~~ electrostatic coupling
on the surface of the substrate can be used.

25 ~~(Synthesis of oligonucleotide~~Oligonucleotide on the
substrate)Substrate

Synthesis of DNA on the substrate includes synthesis on
the silicon substrate using photolithography as a methodology
disclosed in U.S. Patent No. ~~5445934~~. ~~The U.S. Patent No.~~
30 ~~5445934 shows~~5,445,934. This patent discloses a method in
which high density DNA probe arrays are prepared by dividing
the surface of the silicon substrate into very small areas,
and synthesizing DNA for probes. On the other hand, in the
detection substrate for use in the detecting method of the
35 present invention, for example, the size of the section in
which each probe is fixed may be a 0.5 mm square or larger.
Thus, and thus it is not always necessary to enhance the
density. However, also in the detection substrate of the
present invention, photodecomposable protective groups,
40 protective groups that are decomposed by chemicals and the

5 like, are bound to the nucleic acid in advance, and processes of masking, light exposure and reaction are repeated, whereby DNA chains can be synthesized on each section using the methodology described in ~~the U.S. Patent Publication No. 5445934~~ U.S. Patent No. 5,445,934 in which four types of
10 nucleic acid bases are bound for each base to stretch the DNA chain having desired base sequences.

~~(Fixation of oligonucleotide synthesized or collected in advance)~~

15 Fixing Oligonucleotide Synthesized or Collected in Advance

As means for carrying out ~~fixation~~ the fixing using electrostatic coupling, a method in which polylysine, polyethyleneimine and polyalkylamineaone on the solid surface substrate are subjected to blocking using the negative charge
20 of DNA is generally used.

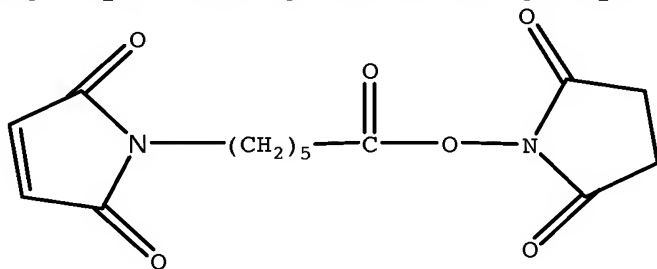
However, in the case of the oligonucleotide with a base length of 60 or less ~~that, which~~ is not sufficiently long, the electric charge of its phosphate groups is also weak. Thus, ~~and thus~~ the binding ~~onto~~ with the substrate by the above -
25 described method is not necessarily strong. For ~~this~~ the oligonucleotide ~~whose~~ with a base length that is not sufficiently long, if a method in which an oligonucleotide with functional groups for a covalent bond introduced in the terminal of nucleic acid is synthesized in advance, the
30 substrate is subjected to surface processing suitable for functional groups, and the above -described functional groups are used to accomplish a covalent bond ~~is~~ are used, stronger binding can be achieved, which is more preferable.

Also, in the case where the oligonucleotide is RNA, the
35 above -described method ~~that, which~~ is used for DNA, may be applied. Alternatively, in the case where the oligonucleotide is a peptide nucleic acid, its nucleic acid part may be used to apply the above -described method ~~that, which~~ is used for DNA.

5 ~~(Types of functional groups)~~ Functional Groups for use in
~~fixation by covalent bond between the solid substrate and~~
~~oligonucleotide)~~ Use in Fixing by Covalent Bond Between the
Solid Substrate and Oligonucleotide

10 When the oligonucleotide is fixed on the solid surface
substrate though a covalent bond, functional groups are
generally introduced in oligonucleotide and the solid surface
substrate, respectively, in advance to carry out the reaction
therebetween. For this combination of functions, a preferable
example is a combination ~~such that~~ in which maleimide groups
15 are introduced in the surface of the substrate and thiol
groups (-SH) are introduced in the oligonucleotide.
Specifically, thiol groups (-SH) are bound to the terminal of
the oligonucleotide while the solid surface is subjected to
processing of forming a coating having maleimide groups, ~~and~~
20 ~~when.~~ When the oligonucleotide is supplied to the solid
surface, the thiol groups (-SH) are made to act on and react
with the maleimide groups to perform ~~fixation~~ the fixing
through the formation of a covalent bond.

25 For introducing maleimide groups in the solid surface,
various kinds of methods may be used, ~~and for.~~ For example,
an aminosilane coupling agent is reacted with a glass
substrate, and then a reagent (EMCS reagent: manufactured by
Dojin Co., Ltd.), including N-(6- maleimidocaproyloxy)
succinimide expressed by the following formula, whereby a
30 coating layer having maleimide groups can be formed.



~~For~~ As another example, a reagent containing succinimidyl
4-(maleimidophenyl)butyrate can be used to react with amino
groups, preferably.

5 Also, for example, an oligonucleotide with thiol groups
introduced therein can be synthesized by using 5' -Thiol-
Modifier C6 (manufactured by Glen Research Co., Ltd.) as a
five prime-end reagent when DNA is synthesized using a DNA
automatic synthesizing apparatus. Furthermore, after
10 synthesis, purification processing by high speed liquid
chromatography is applied after a normal deprotection
reaction.

Combinations of functional groups capable of being used
for ~~fixation~~fixing by the covalent bond include, for example,
15 a combination of epoxy groups (on the solid surface) and amino
groups (the terminal of oligonucleotide) in addition to the
above -described combination of thiol groups and maleimide
groups. Methods for introducing epoxy groups in the solid
surface include, for example, a method in which a coating is
20 applied to the solid surface constituted by polyglycidyl
methacrylate having epoxy groups and a method in which a
silane coupling agent having epoxy groups is applied to the
solid surface made of glass and is reacted with glass.

~~(Supply of oligonucleotide solution)~~Oligonucleotide
25 Solution by the ink-jet process)Ink Jet Process

There is no particular limitation on the means for
supplying predetermined sections on the surface of the solid
substrate with a solution containing the oligonucleotide to be
fixed thereon, as long as a uniform amount of liquid is
30 supplied for each unit area. In the case where printing by
the ink jet process and the like is used, a "solid print
pattern" is prepared. Then, ~~and then~~ using an ink jet type
printer head that is used for ink jet printers, the cartridge
for the ink is filled with an oligonucleotide solution instead
35 of the ink, and printing for a defined area is carried out.
If the amount of the liquid to be supplied is small, items

5 ~~of with a large volume-like, such as an ink cartridge,~~ are not
used. ~~Instead, and instead~~ a structure in which a sample
supplying portion, such as a tube, is connected to a head to
supply the oligonucleotide solution to the head may be used.

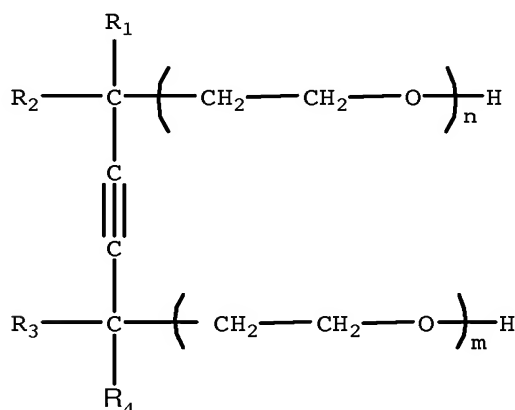
For the oligonucleotide solution for the discharge, which
10 is used in this method, a solution that is capable of being
discharged in the form of ink jets, and has a viscosity
suitable for a minimal amount of droplets discharged from the
head to be shot onto a desired position is used. In addition,
a solvent to be used is selected from solvents that satisfy
15 the above -described requirements and ~~give no damages to~~ do not
damage the desired oligonucleotide in the state of being mixed
with the desired oligonucleotide and during discharge.

Specifically, in terms of dischargeability from the ink
jet head, particularly from the bubble jet head, it is
20 preferable that, for example, the viscosity is in the range of
1 to 15 cps, and the surface tension is 30 dyn/cm or larger,
as the properties of the solution. In particular, when the
viscosity ~~being~~ in the range of 1 to 5 cps and the surface
tension ~~being~~ in the range of 30 to 50 dyn/cm are selected,
25 the position ~~into~~ which the solution is shot onto the
substrate is extremely accurate, and a supplying method using
the bubble jet head is particularly suitably used.

In addition, ~~if~~ in terms of the stability of the
oligonucleotide during discharge and the like ~~are considered,~~
30 the supplying means of the ink jet system is further preferred
when, for example, a solution containing the oligonucleotide
of 2 to 100 mer, particularly ~~of 2 to 60 mer in,~~ at
concentrations ranging from 0.05 to 500 μ M, preferably from 2
to 50 μ M, is used.

35 In applying a discharging method of the ink jet system,
the liquid composition of the oligonucleotide solution is not

5 particularly limited, as long as the solution ~~gives~~
practically ~~no damages to~~ does not damage the desired
oligonucleotide in the state of being mixed with the desired
oligonucleotide and during discharge ~~as a matter of course,~~ as
described above, and it can be discharged to the surface of
10 the solid substrate using the ink jet. Furthermore, a
preferable ~~is a solution containing~~ contains, for example,
glycerin, urea, thiodiglycol or ethylene glycol, isopropyl
alcohol, or acetylene alcohol expressed by the following
formula in addition to desired oligonucleotide:



15
. .
In the above formula, R^1 , R^2 , R^3 and R^4 represent alkyl
groups, for example linear or branched alkyl groups having 1
to 4 carbon atoms, respectively, and m and n represent 0 or
positive integer numbers, respectively, and satisfy $1 \leq m +$
20 ~~$n \leq 30$~~ $n \leq 30$. In addition, specifically, the liquid composition,
including 5 to 10 wt% of urea, 5 to 10 wt% of glycerin, 5 to
10 wt% of thioglycol, and 0.02 to 5 wt%, more preferably 0.5
to 1 wt% of acetylene alcohol expressed by the formula (I),
allows the discharging method of the ink jet system to be ~~used~~
25 suitably used.

(Structure of ~~matrices composed of hydrophobic walls and~~
~~hydrophilic wells~~) Matrices Composed of Hydrophobic Walls and
Hydrophilic Wells

5 Also, for sections of ~~the matrix-form~~ that are provided
on the solid surface, for example, sections of ~~the matrix form~~
~~constituted by~~with hydrophobic walls (barriers) surrounding
hydrophilic wells (recesses) may be formed to prevent coupling
between adjacent sections. A structure may also be used in
10 which the ~~solution of oligonucleotide~~ solution is supplied to
the hydrophilic wells (recesses) surrounded by the hydrophobic
walls (barriers), and the oligonucleotide is fixed only in the
bottom of the hydrophilic wells (recesses).

~~(Materials of walls/wells)~~ Walls/Wells

15 When ~~as~~the sections are arranged in a matrix form, the
solution of oligonucleotide is supplied to the bottom of the
wells (recesses) separated by wall (barrier) patterns to carry
out the binding reaction, ~~it.~~ It is desirable that the bottom
of the wells (recesses) is wetted densely with the solution,
20 but the walls (barriers) have poor wettability with the
solution. For example, it is preferable that the solid
material constituting the surface of the bottom of the wells
(recesses) is ~~much~~more hydrophilic, and the surface of the
walls (barriers) and the portion corresponding to partitions
25 with neighboring sections are less hydrophilic. The ~~solution~~
~~of oligonucleotide~~ solution supplied in the bottom of the well
(recess) is spread across the bottom, but is prevented from
finding its way over the wall (barrier) into adjacent
sections. Also, even the droplet erroneously supplied ~~in~~at
30 the position related to the wall (barrier) quickly moves into
a desired well (recess) having good wettability, ~~and as.~~ As a
result, a predetermined amount of the oligonucleotide solution
can be supplied in the well (recess) more reliably.

35 An example of sections arranged in a matrix form that is
provided on the detection substrate of the present invention
is shown in FIG. 10. The sections in a square matrix form

5 have a structure in which heights (walls) having frame
structures are provided on the surface of the solid substrate,
and arranged rectangular recesses (wells) are separated.
Specifically, the recesses (wells) separated from one another
by the heights (walls) having frame structures are formed by
10 coating the entire surface of the solid substrate with a
material forming heights (walls), and thereafter providing
rectangular through-holes (cut-off portions) to open recesses
(wells). Thus, the bottom of the recess (well) has an exposed
surface of the solid substrate. The exposed portion of the
15 surface of the solid substrate is subjected to processing for
providing a surface to which the oligonucleotide can be bound.
As a result, the oligonucleotide is fixed only in the bottom
of this recess (well).

Materials forming heights (walls) having frame structures
20 include, for example, metals (chrome, aluminum, gold, etc.)
and resins. Resins include ~~resins such as~~, for example,
acryl, polycarbonate, polystyrene, polyimide, acrylate
monomers and urethane acrylate, and photosensitive resins,
such as photoresists, having black dies and black pigments
25 contained therein. Furthermore, for specific examples of
photosensitive resins, UV resists, DEEP-UV resists,
ultraviolet cured resins and the like can be used. UV resists
may include negative resists, such as cyclized polyisoprene-
aromatic ~~pi~~isazidebisazide resists, phenol resin-aromatic azide
30 compound resists, and positive resists, such as novolac resin-
diazonaphthoquinone resists.

DEEP-UV resists may include, for example, radiation
dispersion type polymer resists, such as polymethyl
methacrylate, polymethylene sulfone, polyhexafluorobutyl
35 methacrylate, polymethyl isoprobenil ketone and bromo poly 1-
trimethylcylilpropine, and dissolution inhibiting resists,

5 such as cholate o-nitrobenzyl ester as positive type resists,
and may include borovinylphenol-3-3'- diazidediphenylsulfone,
and polymethacrylate glycidyl as negative type resists.

Ultraviolet cured resins may include polyester acrylate,
epoxy acrylate and urethane diacrylate containing
10 approximately 2 to 10% by weight of one or more types of
photopolymerization initiators, which are selected from
benzophenone and substituted derivatives thereof, oxime
compounds, such as benzyl, and so on.

When detection is carried out using a fluorescent mark, a
15 light-blocking material can be used effectively for curbing a
light reflex by the material forming this height (wall) having
a frame structure. For providing a light-blocking property,
it is effective to add black pigments in the above described
resins, ~~and in.~~ In this case, black pigments that can be used
20 may include carbon black and black organic pigments.

Furthermore, if the height (wall) having a frame
structure is formed by the above described hydrophobic resin,
the surface of the height (wall) is hydrophobic. The
configuration in which heights (walls) having frame structures
25 that are formed by hydrophobic materials are provided is more
preferable ~~in the case where~~ when an aqueous solution is used
as a solution containing the oligonucleotide to be supplied to
the surface of the substrate of recesses (wells). Even if the
aqueous solution is supplied in a position related to the
30 surface of the height (wall), it is not persistently attached
to the surface of the wall, but gradually moves to the bottom
of the recess (well) located in a lower position. Also,
solutions of different oligonucleotides are supplied to
adjacent recesses (wells), but they are separated from each
35 other by the hydrophobic height (wall). Therefore, ~~and~~

5 ~~therefore~~ intermingling (cross -contamination) between solutions due to the penetration of the liquid is prevented.

Furthermore, for the thickness (height from the solid surface) of the height (wall) having a frame structure, the volume of the recess (well) is selected in ~~the light~~ view of
10 the amount of the oligonucleotide solution that is supplied to the recess (well), and the thickness is determined as appropriate so that the volume is filled with the solution. Also, depending on methods of forming the height (wall), the thickness is ~~preferable~~ preferably selected such that it is in
15 the range of 1 to 20 μm and satisfies the above -described requirement. The thickness of the height (wall) selected in this way is in the range of the thickness allowing to effectively prevent cross -contamination between adjacent wells ~~to be prevented effectively~~ when the oligonucleotide
20 solution is supplied to each well by the ink jet process.

~~(Types of specimens)~~ Specimens

Object components contained in the test sample to which the detecting method of the invention can be applied include mRNA, cDNA, proteins, cell extracts and chemicals, such as
25 drugs.

Furthermore, when cDNA is used as an object component, it is possible to use double-strand cDNA directly, ~~but.~~ However the single-strand cDNA marked in advance is preferable in forming hybrid substances efficiently and performing detection
30 thereof conveniently.

On the other hand, mRNA is ~~of~~ essentially single ~~strand~~ stranded, and it is marked in some way to form marked mRNA, thereby making it possible to form hybrid substances efficiently and to perform detection thereof. Furthermore,
35 the amount of mRNA in the test sample is generally small, and it is an object component more remarkably reflecting the

5 advantage that the amount of sample solution required for
detection can be reduced to a low level, which is
characteristic of the detecting method of the present
invention. However, since ~~admixture of~~ RAN decomposition
enzymes ~~tends~~tend to ~~occur~~admix during handling, a
10 predetermined amount of a substance to curb~~prevent~~
decomposition of mRNA, such as RNA decomposition enzyme
inhibitors, such as diethyl pyrocarbonate, is desirable added
in the test sample solution. In addition to mRNA, similarly,
the genome of RNA viruses can be an object component. In
15 addition, tRNA, ribosomal RNA and the like can be object
components.

On the other hand, when the protein is used as an object
component, formed complexes can be detected using the
fluorescence emitted by the protein itself.

20 Also, some chemicals ~~also~~ emit their own fluorescence,
~~and enables~~enabling formed complexes to be detected using the
fluorescence. Chemicals that do not emit fluorescence may be
marked by methods using functional groups of compounds. Those
to which the detecting method of the invention can be applied
25 may include, for example, chemicals that can be bound to
single-strainstranded DNA. In addition, they may include, for
example, chemicals that can be bound to single-strainstranded
RNA.

~~(Means for spotting test samples)~~Spotting Test Samples in
30 ~~an array form)~~Array Form

In the detecting method of the present invention, the
test sample is spotted in an array form in a defined position
on the detection substrate. For the purpose of reducing the
amount of the required liquid to a minimal level, the spot
35 diameter is selected so that it is in the range of several
tens to 100 μm . However, ~~but~~ with such a spot diameter, the

5 liquid should be spotted in high uniformity of spotted amounts and high positional accuracy. As a means for satisfying this requirement, ~~there are~~ spotting apparatuses of pin systems, ink jet systems and capillary systems may be used.

The pin system refers to a method in which the test
10 sample is attached to the pin tip, and the end point thereof is mechanically contacted with the solid surface, thereby taking out a fixed amount of the test sample. The capillary system using capillaries refers to a method in which the test sample solution is sucked up to the capillary on a temporary
15 basis, and the tip of the capillary is mechanically contacted with the solid surface as in the case of the pin system, thereby taking out a fixed amount of the test sample. A ~~various~~Various kinds of spotting apparatuses adopting these two systems are commercially available. Thus, and thus
20 commercially available apparatuses may be used.

The spotting apparatuses of the pin system and capillary system enable any types of test samples to be spotted, and are considered as the most preferable methods for unknown test samples. For example, however, the viscosity of the test
25 sample solution is varied depending on the length and the concentration of DNA contained in the test sample. Therefore, ~~and therefore~~ the amount of the spotted liquid ~~is varied~~varies. Thus, a problem arises in terms of quantification. Also, with respect to proteins, the viscosity
30 of the test sample solution ~~is varied~~varies depending on the size of the molecules and the concentration, thus raising a problem in terms of quantification.

~~(Spots in an array form of test samples by the ink jet process)~~Array Form of Test Samples by the Ink Jet Process

35 Specimens that can be discharged by the ink jet process include chemicals in addition to nucleic acids and proteins.

5 In the ink jet process, because a shearing force is
exerted, the length of nucleic acids and the size of proteins
that can be discharged are limited. However, it is superior
in quantification to the pin system and capillary system, and
is used more suitably than other systems, particularly with
10 respect to the discharge of chemicals. Preferably,
dischargeable nucleic acids are those with a relative length
to bases of 5 kb or smaller, and dischargeable proteins are
those of 1000 K daltons or less. As for chemicals, their
molecular weights are generally small enough compared to
15 nucleic acids and proteins. Therefore, and therefore any
~~chemicals~~ chemical can be discharged, except for polymers
having extremely large molecular weights.

FIG. 3 illustrates schematically a method of discharging
specimen solution by the ink jet process, particularly the
20 bubble jet process, which is one means that is used for
spotting test sample solution in the present invention. In
FIG. 3, reference numeral 101 denotes a liquid supply system
(nozzle) retaining a solution including a specimen as a
discharge liquid in such a manner that the solution is capable
25 of being discharged, reference numeral 103 denotes a solid
phase having a nucleic probe bound thereto with which the
specimen is reacted, and reference numeral 105 denotes a
bubble jet head ~~having a function of giving~~ for providing heat
energy to the liquid to discharge it, which is a type of ink
30 jet head. Reference numeral 104 denotes a liquid (droplet)
including the specimen discharged from the bubble jet head.
FIG. 4 is a sectional view of the bubble jet head 105
described in FIG. 3. In FIG. 4, reference numeral 107 denotes
a liquid including a specimen solution to be discharged from
35 the bubble jet head 105, and reference numeral 118 denotes a
substrate portion having a heat generation portion to

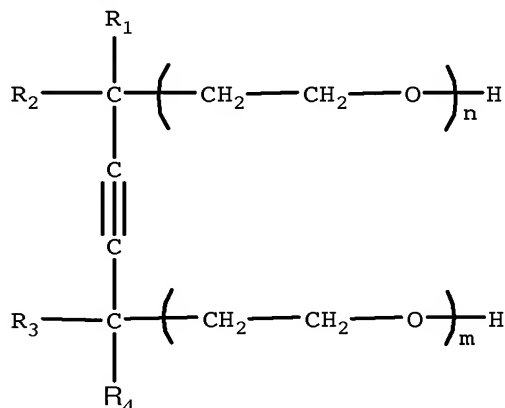
5 ~~give~~provide discharge energy to the above -described liquid.
The substrate portion 118 includes a protective layer 109
formed by silicon oxide and the like, electrodes 111-1 and
111-2 formed by aluminum and the like, an exothermic resistor
layer 113 formed by nichrome and the like, a heat storage
10 layer 115, and a support 116 formed by aluminum having good
heat-release ~~property~~properties. The liquid 107 including the
specimen comes to a discharge orifice (discharge outlet)
~~119, 119~~ and forms a meniscus 121 with a predetermined
pressure. In this situation, when ~~electric~~electrical signals
15 are applied to the electrodes 111-1 and 111-2, a region
(foaming region) denoted by reference numeral 123 abruptly
releases heat, and the liquid 117 contacted therewith is
discharged and flies toward the solid surface 103. The amount
of liquid that can be discharged using a bubble jet head
20 having such a structure varies depending on the size of its
nozzle, but can be controlled to approximately ~~to~~ 4 to 50
picoliters, which is extremely useful as a means for arranging
probes ~~in~~at a high density in a matrix form on the surface of
the substrate.

25 ~~And, in~~In terms of dischargeability from the ink jet,
particularly from the bubble jet head, for the properties of
the above -described liquid, it is preferable that its
viscosity ~~is~~be in the range of 1 to 15 cps and its surface
tension ~~is~~be 30 dyn/cm or larger. Also, if the viscosity is
30 in the range of 1 to 5 cps and the surface tension is in the
range of 30 to 50 dyn/cm, the position in which the droplet is
spotted (spot position) on the solid phase is extremely
accurate, ~~allowing the method to be used particularly~~
~~suitably~~.

35 In addition, if the stability of nucleic acid during
discharge or the like is taken into consideration, a single-

~~strain~~stranded nucleic acid or double-~~strain~~stranded nucleic acid of, for example, 2 to 5000 mer, particularly 2 to 10000 mer, is preferably contained in the solution. For example, c-DNA chips are preferably contained ~~in~~at the concentration of 0.05 to 500 μM , particularly 2 to 50 μM .

~~For the composition of discharged liquid, the~~The composition of liquid is not particularly limited, as long as the liquid has no substantial influence on the nucleic acid probe when it is mixed with the nucleic acid probe and when it is discharged from the ink jet, ~~and it.~~ The liquid can be normally discharged to the solid phase using the ink jet, ~~but preferable.~~ Preferable are liquids including glycerin, urea, thiodiglycol or ethylene glycol, isopropyl alcohol, and acetyl alcohol expressed by the following formula:



In the above formula, R^1 , R^2 , R^3 and R^4 represent alkyl groups, specifically linear or branched alkyl groups having 1 to 4 carbon atoms, m and n represent 0 or positive integer numbers, respectively, and satisfy $1 \leq m + n \leq 30$.

Further, specifically, a liquid containing 5 to 10% by weight (wt%) of urea, 5 to 10 wt% of glycerin, 5 to 10 wt% of thiodiglycol, and 0.02 to 5 wt%, more preferably 0.5 to 1 wt% of acetylene alcohol, is suitably used.

{Examples}

5 The present invention will be described in detail below
using Examples. Furthermore, the Examples shown herein
represent ~~one example of~~ most suitable embodiments of the
present invention, ~~but~~and the invention should not be limited
by these Examples.

10 Example 1

A glass substrate with black matrices for specimen
matrices for analyzing sequences of p 53 genes on a specimen
matrix substrate partitioned by patterns is prepared.

1. Preparation of a black matrix introduction substrate
15 coated with polylysine.

A glass substrate (60 mm x 50 mm) made of synthetic
quartz is subjected to supersonic cleaning using 2% sodium
hydrate solution, and is then subjected to UV ozone processing
to clean the surface. Then, a polylysine solution
20 (manufactured by ~~sigma~~Sigma Co., Ltd.) is applied to the
entire surface with a spin coater. In addition, a DEEP-UV
resist (negative type resist for black matrices) (BK-739P
manufactured by Nippon Steel Chemical Co., Ltd.) is applied
thereto with the spin coater so that the thickness after
25 curing is 5 μ m, and this substrate is heated for curing at
80°C for 5 minutes with a hotplate. Using a DEEP-UV aligner,
a region of 1 cm x 1 cm is proximately exposed to light using
a patterned mask, so that the distance (X) between adjacent
wells in FIG. 1 is 100 μ m and the form of the well is a ~~square~~
30 ~~of~~ 1 mm x 1 mm square, and then development is carried out
with a developing ~~solution of~~ inorganic alkaline solution
using a spin drier, and the developing solution is washed out
completely with purified water.

Then, the substrate is briefly dried using the spin
35 drier, and is thereafter heated at 180°C for 30 minutes in a
clean oven to have the resist fully cured to obtain a

5 substrate in which 400 wells are ~~arranged as~~at a predetermined arrangement and adjacent wells are separated from each other by the black matrix. Furthermore, the volume of each well is calculated as 5 μ l if the thickness of the liquid is 5 μ m.

2. ~~Fixation of specimen DNA~~Fixing Specimen DNA

10 (1) Preparation of cDNA libraries

The p 53 gene is obtained by a PCR reaction from 64 types of cDNA libraries obtained from tumor tissues.

That is, RNA samples were obtained from each tissue collected with biopsies using Catrimox-14 (Biotechnology Co.,
15 Ltd.). Based on this sample solution, First-Strand cDNA Synthesis Kit (manufactured by Life Sciences Co., Ltd) is used to obtain cDNA libraries.

(2) Amplification of p53 genes having T3 binding sites by a PCR method.

20 Based on the cDNA library, "Human p53 Amplimer set" manufactured by CLONTECH Co., Ltd. is used to carry out the PCR reaction.

As a PCR reaction solution, "one shot LA PCR Mix" (Takara Shuzo Co., Ltd.) was used. The composition of the PCR
25 reaction solution is as follows:

One shot LA PCR Mix	25 μ l
5' primer (20 μ M)	1
3' primer (20 μ M)	1
cDNA library solution	1
DW	22/50 μ l.

The PCR cycle is such that after thermal denaturation at 95°C for 5 minutes, cycles at 95°C for 30 seconds, at 55°C for 30 seconds and at 72°C for 60 seconds are conducted 29 times.
Finally, ~~and finally~~ the solution is left for a reaction at
30 72°C for 5 minutes and is then stored at 4°C.

5 After the reaction, gel electrophoresis is performed to confirm a product existing in the region of molecular weight of about 300 mer, and purification is carried out with MicroSpin Column S200 (Pharmacia) to obtain p 53 genes (p 53 DNA).

10 (3) Synthesis of single-~~strain~~stranded p 53 DNA

Using as a matrix the DNA obtained in the above (2), a single-~~strain~~stranded marked DNA is obtained by the PCR reaction using 5' primer (Takara Shuzo Co., Ltd.). The composition of the reaction solution comprises

One shot LA PCR Mix	25 μ l
5' primer (20 μ M)	1
P 53 DNA	1
DW	22/50 μ l, and

15 the reaction cycle is such that cycles at 96°C for 30 seconds, at 60°C for 15 seconds and at 60°C for 4 minutes are repeated 24 times, and finally, the solution is stored at 4°C. Thereafter, it is purified with MicroSpin Column S200.

(4) ~~Fixation of~~Fixing p 53 cDNA

20 5 μ l of the single-~~strain~~stranded DNA obtained in the above (3) is injected under a microscope into each well of the polylysine-coated substrate with black matrices prepared in the above (1), and is fixed through electrostatic coupling.

3. Analysis of variation of p 53 genes with
25 oligonucleotide probes

The 64 DNAs were selected, focusing the attention on the 248th and 249th amino acid sequences of the p 53 gene being a tumor inhibitor gene. That is, it is known that a case of frequent variation in the base sequence of CGGAGG is the case
30 where the first C is changed to T, the second A is changed to G, and the third G of the sequence corresponding to the 249th

5 amino acid is changed to T. Thus, the 64 probes are designed,
focusing the attention on the base sequence at these three
points.

That is, it is a structure in which the total length of
the probe is 18 mer, and six bases including this variation
10 are located at the center, thereof, and the bases are
sandwiched between common sequences. A common sequence
corresponds to the range from the five prime-end to the
ATGAAC, and the subsequent portion including variation
corresponds to the NNGAGN and a further subsequent common
15 portion corresponds to the CCCATC, resulting in a final
sequence of 5'ATGAACNNGAGNCCCATC3' (SEQ ID NO:65). Here, the
portion expressed by N corresponds to the A, G, C, and T that
are four types of nucleic acid bases. The probe DNA has a
sequence complementary to the sequence to be detected (the
20 above described sequence), and thus the sequence thereof is
~~5'MCGGNCTCNGTTTCAT3' (SEQ ID NO:66).~~ 5'GATGGGNCTCNGTTTCAT3'
(SEQ ID NO:66). Rhodamine is coupled to the five prime-end of
each probe sequence to mark the probe. Specific base
sequences of these 64 types of marked DNA probes are shown in
25 the following Table 1.

5

Table 1

SEQ ID NO.	Sequence	SEQ ID NO.	Sequence
1	GATGGGACTCAAGTTCAT	33	GATGGGCCTCAAGTTCAT
2	GATGGGACTCAGGTTCAT	34	GATGGGGCTCAGGTTCAT
3	GATGGGACTCACGTTCAT	35	GATGGGCCTCACGTTCAT
4	GATGGGACTCATGTTCAT	36	GATGGGCCTCATGTTCAT
5	GATGGGACTCGAGTTCAT	37	GATGGGCCTCGAGTTCAT
6	GATGGGACTCGGGTTCAT	38	GATGGGCCTCGGGTTCAT
7	GATGGGACTCGCGTTCAT	39	GATGGGCCTCGCGTTCAT
8	GATGGGACTCGTGTTTCAT	40	GATGGGCCTCGTGTTTCAT
9	GATGGGACTCCAGTTCAT	41	GATGGGCCTCCAGTTCAT
10	GATGGGACTCCGGTTCAT	42	GATGGGCCTCCGGTTCAT
11	GATGGGACTCCCGTTCAT	43	GATGGGCCTCCCGTTCAT
12	GATGGGACTCCTGTTCAT	44	GATGGGCCTCCTGTTCAT
13	GATGGGACTCTAGTTCAT	45	GATGGGCCTCTAGTTCAT
14	GATGGGACTCTGGTTCAT	46	GATGGGCCTCTGGTTCAT
15	GATGGGACTCTCGTTCAT	47	GATGGGCCTCTCGTTCAT
16	GATGGGACTCTTGTTTCAT	48	GATGGGCCTCTTGTTTCAT
17	GATGGGGCTCAAGTTCAT	49	GATGGGTCTCAAGTTCAT
18	GATGGGGCTCAGGTTCAT	50	GATGGGTCTTAGGTTCAT
19	GATGGGGCTCACGTTCAT	51	GATGGGTCTCACGTTCAT
20	GATGGGGCTCATGTTCAT	52	GATGGGTCTCATGTTCAT
21	GATGGGGCTCGAGTTCAT	53	GATGGGTCTCGAGTTCAT
22	GATGGGGCTCGGGTTCAT	54	GATGGGTCTCGGGTTCAT
23	GATGGGGCTCGCGTTCAT	55	GATGGGTCTCGCGTTCAT
24	GATGGGGCTCGTGTTTCAT	56	GATGGGTCTCGTGTTTCAT
25	GATGGGGCTCCAGTTCAT	57	GATGGGTCTCCAGTTCAT
26	GATGGGGCTCCGGTTCAT	58	GATGGGTCTCCGGTTCAT
27	GATGGGGCTCCCGTTCAT	59	GATGGGTCTCCCGTTCAT
28	GATGGGGCTCCTGTTCAT	60	GATGGGTCTCCTGTTCAT
29	GATGGGGCTCTAGTTCAT	61	GATGGGTCTCTAGTTCAT
30	GATGGGGCTCTGGTTCAT	62	GATGGGTCTCTGGTTCAT
31	GATGGGGCTCTCGTTCAT	63	GATGGGTCTCTCGTTCAT
32	GATGGGGCTCTTGTTTCAT	64	GATGGGTCTCTTGTTTCAT

5 Then, for each of the 64 types of marked probe DNAs, a 8
µM solution containing glycerin, urea and thiodiglycol ~~in at~~
the final concentration of 7.5%, and acetylenol EH ~~in at~~ the
final concentration of 1% is prepared. A different probe
solution is charged by 100 µl in each of the six nozzles of BJ
10 Printer Head BC 62 (manufactured by ~~Cannon~~Canon Inc.).
~~Arrangement~~An arrangement is made so that six DNAs can be
discharged for each head, and two heads are used to discharge
12 DNAs at a time, and the heads are exchanged 6 times to
discharge DNAs so that each spot of 64 DNA is formed
15 independently. In this ~~way,~~manner, a total of 64 probes are
discharged in the form of the 8 x 8 array in each well of a
black matrix coated with polylysine.

FIG. 5 shows an arrangement on each black matrix of 64
DNA probes that are discharged. In this case, 64 DNA probes
20 are spotted in one matrix.

Thereafter, this substrate in which each probe is spotted
is left in a humidifier chamber set at 40°C to carry out a
hybridization reaction.

Thereafter, the substrate is cleaned with a 10 mM
25 phosphate buffer containing 100 mM NaCl to remove DNA probes
that have not been engaged in the formation of the hybrid
substance.

DNA arrays after the hybridization reaction are observed
using an inverted fluorescence microscope equipped with a
30 filter set ~~suitable~~suitably for rhodamine.

If the gene as a specimen has normal base sequences,
spots of highest fluorescence intensity should be observed in
the gene at the location of the relative 42~~th~~end DNA probe. It
can be considered that those are derived from the hybrid of
35 the p 53 gene having normal sequences ~~amplified with~~exposed to

5 the probe DNA and PCR. In a varied gene, detectable spots are observed at the location other than the 42~~th~~nd location, and a varied sequence can be ~~known~~identified from the DNA probe supplied to the location.

Example 2

10 ~~(Evaluation of existence/not-Existence/Non-existence of carcinogenic genes using~~Carcinogenic Genes Using mRNA)

1. Extraction of mRNA

"QuickPrep Micro mRNA Purification Kit" (manufactured by Amersham Pharmacia ~~biotech~~Biotech co., Ltd.) is used to
15 extract mRNA from tumor tissues collected with the biopsy. This mRNA is bound to a polylysine substrate with black matrices, as in the case of Example 1.

2. Examination of existence/~~not-non~~-non-existence of carcinogenic genes and the type thereof with various kinds of
20 carcinogenic gene probe arrays

Sets of cloned oncogenes (18 types, manufactured by Takara Shuzo Co., Ltd.) are purchased. Then, and then "LabelITnon-RI Labeling Kits" are used to perform rhodamine marking.

25 18 types of marked oncogene probes are spotted as an arrangement of 4 x 5 on the above -described substrate with mRNA bound thereto, using a microarray preparing apparatus (pin system) manufactured by Cartesian Technologies Co., Ltd.

Further, a hybridization reaction is carried out as in
30 ~~the case of~~ Example 1.

The type of oncogenes existing in the mRNA section extracted from each tissue can be known.

At this time, sufficient detection can be ~~sufficiently~~ performed with one type of marks irrespective of the types of
35 oncogenes present.

5 The second aspect of the invention ~~will be describe~~is
described more specifically below with reference to the
Examples ~~below~~.

~~(Example 3)~~

 An example of procedures for preparing ~~ana~~ substrate with
10 oligonucleotide bound thereto will be described below. In
this embodiment, a detection substrate with the
oligonucleotide bound to a region of 2 mm square on a glass
substrate was prepared in accordance with the procedure
described below.

15 1. Cleaning of the substrate

 A glass substrate of 1-inch square was placed ~~in on~~ a
rack, ~~and was~~ soaked in a detergent for ultrasonic cleaning.
Thereafter, it was subjected to ultrasonic cleaning in the
above _described detergent for 20 minutes, followed by
20 removing the detergent by rinsing. ~~Furthermore~~Also, it was
rinsed with distilled water, followed by further performing
ultrasonication for 20 minutes in a container containing
distilled water.

 Then, this glass substrate was soaked for 10 minutes in
25 1N sodium hydrate solution heated in advance. After it was
taken out from the solution, the 1N sodium hydrate solution
adhered to the surface was washed out with water. Thereafter,
~~and thereafter~~ cleaning with distilled water was continued.

 2. Surface treatment

30 The above _described cleaned glass substrate was soaked
in an aqueous solution of 1% silane coupling agent
(manufactured by Shin-Etsu Chemical Co., Ltd., Trade name:
KBM 603) at room temperature for 20 minutes, followed by
spraying nitrogen gas on ~~the~~ both sides of the substrate to
35 drive off water for drying. The substrate was baked for one
hour by using an oven heated to 120°C to complete the

5 treatment of the surface of the glass substrate with a silane coupling agent.

On the other hand, 2.7 mg of EMCS (N-(6-Maleimidocaproyloxy) succinimide: manufactured by Dojin Co., Ltd.) was weighed, and was dissolved in a solution of
10 DMSO/ethanol (1:1) (final concentration of 0.3 mg/ml). The glass substrate subjected to the treatment with a silane coupling agent was soaked in this EMCS solution for two hours to carry out the reaction between the amino group of the silane coupling agent covering the surface of the substrate
15 and the succinimide group in the EMCS solution. In association with this reaction, the substrate is covered with EMCS through the silane coupling agent. In the obtained glass surface, a maleimide group derived from the EMCS ~~exists~~is present on the surface. The glass substrate taken out after
20 the reaction with the EMCS solution is cleaned with distilled water, and is thereafter dried with nitrogen gas. This glass substrate subjected to the surface treatment for introducing a maleimide group will be used for a binding reaction with DNA described ~~later~~below.

25 3. Synthesis of DNA for fixing glass substrates

~~Oligonucleotide~~An oligonucleotide having a base sequence of the following Sequence 1 (SEQ ID NO:42) is chemically synthesized for ~~fixation~~fixing on the glass substrate. This sequence 1 is a 18 mer sequence including in its central part
30 a base sequence with a base length of 6 to code 248th and 249th amino acids in an amino acid sequence of a gene product (peptide chain) that is coded by the p 53 gene known as a tumor suppressor gene. Also, A SH group is introduced in its 5' end for fixation on the glass substrate.

35 Sequence 1 5' HS-GATGGGCCTCCGGTTCAT3' (SEQ ID NO:42)

5 The SH group is introduced by using a commercially
available reagent Thiol-Modifier (manufactured by GlenResearch
Co., Ltd.) on a DNA automatic synthesizing apparatus.
Subsequently, normal ~~deprotection~~deprotection was carried out
to recover DNA, ~~and the~~. The DNA was purified by high -speed
10 liquid chromatography, and was then used in the following
processes.

4. Discharging of DNA using a BJ printer head and
binding thereof to a substrate

The above -described synthetic oligonucleotide (DNA) was
15 dissolved in water, and the solution was diluted to ~~the~~ a
concentration of 8 μ M using SG Clear (a solution containing
7.5% of glycerin, 7.5% of urea, 7.5% of thiodiglycol and 1% of
acetylenol EH).

100 μ l of this oligonucleotide solution was charged into
20 the nozzle of BJ printer head BC 62 (manufactured by Canon
Inc.) with the nozzle modified so that it is suitable for a
small amount of samples (discharged amount). This modified
printer head was set in a plotting apparatus to perform
printing over the surface of the glass substrate as an area of
25 "solid print" of 2 mm square with the oligonucleotide
solution. Furthermore, the modified printer head that was
used is used for bubble jet type ink jet printing and enables
printing to be performed at a resolution of 360 x 720 dpi.

Thereafter, the glass substrate coated with the
30 oligonucleotide solution was left in a humidifier chamber for
30 minutes to carry out a reaction between the maleimide group
on the surface of the substrate and the thiol group (HS-) of
the oligonucleotide. Thereafter, the unreacted
oligonucleotide was removed. The prepared substrate ~~to detect~~
35 ~~is a substrate with~~for detection has the synthetic DNA

5 (oligonucleotide) of the above -described Sequence 1 bound to a predetermined ~~section of~~ 2 mm square section on the glass substrate ~~through~~via a covalent bond.

(Example 4)

Supply of cDNA solution to the ~~surface of the substrate~~
10 ~~with oligonucleotide bound thereto and hybridization~~
~~reaction.~~Surface of the Substrate with Oligonucleotide Bound
Thereto and Hybridization Reaction

From various kinds of cDNA libraries obtained from tumor tissues, p 53 gene fragments were PCR-amplified, and then only
15 one type of side chains ~~were~~was reamplified using primers marked in advance to prepare marked single-~~strain~~stranded cDNA for use as test samples. The hybridization reaction was carried out between this marked single ~~strain~~stranded DNA derived from the p 53 gene and the DNA probe bound on the
20 detection substrate prepared~~7~~ in Example 3.

1. Preparation of test samples

From 64 types of cDNA libraries obtained from tumor tissues, p 53 gene fragments were obtained by the PCR reaction.

25 Specifically, first, all RNA samples were separated/collected from respective tissues collected ~~with~~
~~the~~via a biopsy, using Catrimox-14 (Biotechnology Co., Ltd.). On the basis of ~~the~~all the RNA sample solutions, a c-DNA library was prepared using First-Strand cDNA Synthesis Kit
30 (manufactured by Life Science Co., Ltd.). A primer for amplifying p 53 genes was added to this cDNA library to amplify P 53 gene fragments. With this PCR amplification product as a template, the marked five -side primer was used to carry out the PCR reaction (DNA synthetic reaction) to
35 amplify only one type of side chains. By this amplification,

5 marked single ~~strain~~stranded DNA derived from the p 53 gene
can be prepared.

(1) Amplification of p 53 gene fragments having a T3
binding site in the terminal by the PCR method.

For using a primer for auto sequencers (Takara Shuzo Co.,
10 Ltd) using T3 promoters as the above described marked primer,
a primer having a T3 site in the terminal and having coupled
to its downstream a base sequence allowing the p 53 gene part
to be amplified was first synthesized. The PCR reaction was
carried out using this primer to obtain a PCR amplification
15 product having a T3 promoter site coupled to the p 53 gene
part.

In this example, for the five prime-end primer for
amplifying p 53 genes, the primer with a base sequence having
a T3 promoter site coupled to its five side (T3-P53)was
20 prepared. The base sequence is shown below.

5'

AATTAACCCTCACTAAAGGGAACCTGAGGTTGGCTCTGACTGTACCACCATCC3' (SEQ
ID NO:67)

In the sequence, the underlined part on the side of five
25 prime-end represents a T3 polymerase binding site. On the
other hand, for a three prime-end primer for amplification, a
three prime-end primer attached in a commercially available
amplification kit, "Human p 53 Amplimer Set" of CLONTECH Co.,
Ltd.was used. For a PCR reactive solution, "one shot LA PCR
30 Mix" (Takara Shuzo Co., Ltd.) was used.

The solution composition in the PCR reaction has:

one shot LA PCR Mix	25 μ l
T3-P53 primer (20 μ M)	1 μ l
3' primer (20 μ M)	1 μ l
cDNA library solution	1 μ l

DW

22 μ l/50 μ l ~~7.~~

5 ~~and for~~For the PCR cycle, ~~a condition of conducting the~~
cycles were conducted at 95°C for 30 seconds, at 55°C for 30
seconds and at 72°C for 60 seconds at 29 times after thermal
denaturation at 95°C for 5 minutes, and finally keeping the
solution at 72°C for five minutes ~~was used, and the.~~ The
10 reactant was stored at 4°C on a temporary basis after it was
cooled.

After the reaction, gel electrophoresis was carried out
to confirm a PCR product ~~existing~~present in the region of
molecular weight of about 300 mer. This PCR product was
15 purified with Micro Spin Column 5200 (Pharmacia) to obtain p
53 gene fragments to which the T3 primer can be coupled (T3-
linked p 53 DNA).

(2) Synthesis of marked single ~~strain~~stranded DNA using
labeled T3 primers (Rho-T3) ~~+~~

20 With the p 53 gene fragment obtained in (1) as a matrix,
single ~~strain~~stranded marked DNA was obtained with the PCR
reaction, using a Rho-T3 primer (Takara Shuzo Co., Ltd.).
The composition of the reactive solution had:

one shot LA PCR Mix 25 μ l

Rho-T3 primer (10 μ M) 1 μ l

T3-linked p 53 DNA 1 μ l

DW

23 μ l/50 μ l ~~7.~~

~~and for~~For the reaction cycle, ~~a condition of~~
25 ~~conducting the~~ cycles were conducted at 96°C for 30 seconds, at
50°C for 15 seconds and at 60°C for 4 minutes 24 times ~~was~~
~~used, and the.~~ The reactant was stored at 4°C on a temporary
basis after it was cooled. It was purified with Micro Spin
Column S200, ~~and thereafter~~200. Thereafter, gel
30 electrophoresis was carried out to confirm desired rhodamine

5 labeled single-~~strain-stranded~~ DNA synthesized through the PCR reaction.

2. Supply of test sample solution

Sodium chloride was added in the test sample obtained in the above described process, namely the solution of rhodamine marked single-~~strain-stranded~~ DNA derived from the p 53 gene, so that the final concentration of the solution was 1M. The solution of rhodamine marked single-~~strain-stranded~~ DNA derived from the p 53 gene, which had been prepared from 64 types of c DNA libraries, was injected into each well of a 96-
15 hole microtiter plate. These solutions of rhodamine marked single-~~strain-stranded~~ DNA were spotted ~~as in~~ an 8 x 8 arrangement ~~of 8 x 8~~ onto the detection glass substrate with the DNA probe of Sequence 1 obtained in Example 3 in the form of 2 mm square, using a microarray preparing apparatus (pin system) manufactured by Cartesian Technologies. The diameter
20 of each spot was 100 μ m.

3. Hybridization reaction-

This detection substrate with a total of 64 types of rhodamine marked single-~~strain-stranded~~ DNA solutions being
25 sample specimens spotted thereon was left in a humidifier chamber set at 40°C to carry out a hybridization reaction for 3 hours. Thereafter, the detection substrate was washed with a 10 mM phosphate buffer containing 100 mM NaCl to remove test samples that had not been engaged in the formation of hybrid
30 substances.

After the hybridization reaction, the test sample spotted in the form of a two-dimensional ~~array of 8 x 8~~ array was observed using an inverted fluorescence microscope equipped with a filter set for excitation light and fluorescence
35 suitable for fluorescence marked rhodamine. For ~~the most part~~

5 of the spots, red fluorescence derived from fluorescence
marked rhodamine in association with the formation of hybrid
substances was observed. However, fluorescence intensity was
weak for six spots and no fluorescence was observed for one
spot.

10 ~~For~~Due to this, it can be considered that since in the p
53 gene derived from corresponding six types of tumor cells,
variation occurs somewhere in the base sequence corresponding
to the 248th and 249th of the amino acid sequence of the p 53
gene product (p 53 protein), the amount of formed hybrid
15 substances is small due to its mismatch,~~and in association~~
~~therewith.~~ In that connection, the fluorescence intensity
from the fluorescence mark is weak. For the test sample in
which fluorescence was not observed, it can be considered ~~from~~
~~the fact that~~ because hybrid substances ~~were not formed that~~
20 in p 53 cDNA fragments ~~contained in the sample,~~ did not form, a
deficiency occurs in the base sequence to code the above =
described 248th and 249th ~~of the~~ amino acid sequence.
Consequently, ~~and consequently~~ hybrid substances could not be
formed.

25 ~~(Example 5)~~

Preparation of ~~array form spots of test samples on the~~
~~probe matrix detection substrate with multiple~~
~~oligonucleotides fixed thereon.~~ Array form Spots of Test
Samples on the Probe Matrix Detection Substrate with Multiple
30 Oligonucleotides Fixed Thereon

1. Preparation of 64 probe matrices-

Processing was performed as in ~~the case of~~ Example 3 to
prepare a glass substrate having a maleimide group . 64 DNA
~~of which~~ DNA's with the base sequences are as shown in Table 2
35 were printed (applied) thereon in the area of 2 mm square,
respectively, using a bubble jet printer head similar to that

5 ~~of~~in Example 3 to prepare a detection substrate on which
sections with 64 types of probe DNAs fixed therein were
arranged in a matrix form.

Focusing the attention on the 248th and 249th amino acids
of the amino acid sequence of the gene product (p 53 protein)
10 of the p 53 gene being a tumor suppressor gene, 64 DNAs ~~of~~
~~which~~with base sequences ~~areas~~ shown in Table 1 were selected
on the basis of the base sequence to code these two amino
acids so that a sequence with various kinds of base variations
added thereto was obtained. Specifically, it is known that a
15 case of frequent variation in the base sequence CGGAGG
providing a base is the case where the first C of the CGG to
code the 248th amino acid is changed to T, the second A is
changed to G, and the third G of the AGG to code the 248th
amino acid is changed to T. Thus, 64 probes were designed to
20 provide sequences capable of being bound to base sequences
with these bases at three positions varied in various ~~kinds of~~
~~forms~~ways.

~~Actually~~Specifically, it was a structure in which the
total length of the probe was 18 mer, six bases including this
25 variation were located in the center thereof, and common base
sequences with base lengths of 6 were placed before and after
the six bases. More specifically, the structure has a common
sequence of ATGAAC from the side of the five prime-end, the
base sequence of NNGAGN as a portion including the variation,
30 and a common sequence of CCCATC on the side of three prime-
end.

It was a base sequence ~~a base sequence~~ complimentary to
the sequence of 5'ATGAACNNGAGNCCCATC3' (SEQ ID NO:65). That
is, it was a probe expressed by 5'GATGGGNCTCNGTTTCAT3' (SEQ ID
35 NO:66). Furthermore, since it is a DNA probe, the portion

5 denoted by N in the above _described base sequence refers to any one of A, G, C and T that are four DNA nucleic acid bases.

5

Table 2

1	5'-GATGGGACTCAAGTTCAT-3'	33	5'-GATGGGCCTCAAGTTCAT-3'
2	5'-GATGGGACTCAGGTTCAT-3'	34	5'-GATGGGCCTCAGGTTCAT-3'
3	5'-GATGGGACTCACGTTCAT-3'	35	5'-GATGGGCCTCACGTTCAT-3'
4	5'-GATGGGACTCATGTTCAT-3'	36	5'-GATGGGCCTCATGTTCAT-3'
5	5'-GATGGGACTCGAGTTCAT-3'	37	5'-GATGGGCCTCGAGTTCAT-3'
6	5'-GATGGGACTCGGGTTCAT-3'	38	5'-GATGGGCCTCGGGTTCAT-3'
7	5'-GATGGGACTCGCGTTCAT-3'	39	5'-GATGGGCCTCGCGTTCAT-3'
8	5'-GATGGGACTCGTGTTTCAT-3'	40	5'-GATGGGCCTCGTGTTTCAT-3'
9	5'-GATGGGACTCCAGTTCAT-3'	41	5'-GATGGGCCTCCAGTTCAT-3'
10	5'-GATGGGACTCCGGTTCAT-3'	42	5'-GATGGGCCTCCGGTTCAT-3'
11	5'-GATGGGACTCCCGTTCAT-3'	43	5'-GATGGGCCTCCCGTTCAT-3'
12	5'-GATGGGACTCCTGTTCAT-3'	44	5'-GATGGGCCTCCTGTTCAT-3'
13	5'-GATGGGACTCTAGTTCAT-3'	45	5'-GATGGGCCTCTAGTTCAT-3'
14	5'-GATGGGACTCTGGTTCAT-3'	46	5'-GATGGGCCTCTGGTTCAT-3'
15	5'-GATGGGACTCTCGTTCAT-3'	47	5'-GATGGGCCTCTCGTTCAT-3'
16	5'-GATGGGACTCTTGTTTCAT-3'	48	5'-GATGGGCCTCTTGTTTCAT-3'
17	5'-GATGGGGCTCAAGTTCAT-3'	49	5'-GATGGGTCTCAAGTTCAT-3'
18	5'-GATGGGGCTCAGGTTCAT-3'	50	5'-GATGGGTCTCAGGTTCAT-3'
19	5'-GATGGGGCTCACGTTCAT-3'	51	5'-GATGGGTCTCACGTTCAT-3'
20	5'-GATGGGGCTCATGTTCAT-3'	52	5'-GATGGGTCTCATGTTCAT-3'
21	5'-GATGGGGCTCGAGTTCAT-3'	53	5'-GATGGGTCTCGAGTTCAT-3'
22	5'-GATGGGGCTCGGGTTCAT-3'	54	5'-GATGGGTCTCGGGTTCAT-3'
23	5'-GATGGGGCTCGCGTTCAT-3'	55	5'-GATGGGTCTCGCGTTCAT-3'
24	5'-GATGGGGCTCGTGTTTCAT-3'	56	5'-GATGGGTCTCGTGTTTCAT-3'
25	5'-GATGGGGCTCCAGTTCAT-3'	57	5'-GATGGGTCTCCAGTTCAT-3'
26	5'-GATGGGGCTCCGGTTCAT-3'	58	5'-GATGGGTCTCCGGTTCAT-3'
27	5'-GATGGGGCTCCCGTTCAT-3'	59	5'-GATGGGTCTCCCGTTCAT-3'
28	5'-GATGGGGCTCCTGTTCAT-3'	60	5'-GATGGGTCTCCTGTTCAT-3'
29	5'-GATGGGGCTCTAGTTCAT-3'	61	5'-GATGGGTCTCTAGTTCAT-3'
30	5'-GATGGGGCTCTGGTTCAT-3'	62	5'-GATGGGTCTCTGGTTCAT-3'
31	5'-GATGGGGCTCTCGTTCAT-3'	63	5'-GATGGGTCTCTCGTTCAT-3'
32	5'-GATGGGGCTCTTGTTTCAT-3'	64	5'-GATGGGTCTCTTGTTTCAT-3'

Then, for each of the 64 types of labeled probe DNAs, ~~a~~an 8 μ M solution containing glycerin, urea and thiodiglycol ~~in~~at the final concentration of 7.5%, respectively, and acetylenol EH ~~in~~at the final concentration of 1% was prepared. As in ~~the~~

5 ~~case of~~ Example 4, using BJ Printer Head BC 62 (manufactured
by ~~Cannon~~Canon Inc), a different DNA probe solution was
charged by 100 µl in each of the six nozzles of the printer
head, ~~and using.~~ Using a plurality of such printer heads, a
detection substrate with total 64 DNA probes applied to and
10 fixed in each section of 2 mm square in the form of a "solid
print" and arranged in a matrix form (8 x 8) was prepared. A
schematic layout of the 64 DNA probes arranged in a matrix
form (8 x 8) on the detection substrate is shown in FIG. 7.

2. Preparation of array spots of test samples.

15 As in the case of Example 4, 64 types of labeled cDNAs
were spotted in the form of the two -dimensional 8 x 8 array
on each region of 2 mm square for fixing probes.
Specifically, as schematically shown in FIG. 8, a pin system
array preparing apparatus was used to form spots in the form
20 of the two -dimensional 8 x 8 array on the sections arranged
in a matrix form (8 x 8) in which each DNA probe was fixed.

3. Hybridization reaction-

A hybridization reaction was carried out using conditions
and procedures similar to those ~~of~~in Example 4. The result
25 thereof is shown in FIG. 9. In the arrangement shown in FIG.
7, with respect to spots on probes corresponding to the base
sequence of the 42nd normal gene, fluorescence intensity was
weak for six spots as in ~~the case of~~ Example 4. Also, no
fluorescence was observed for one spot. In addition thereto,
30 it was observed that ~~fluorescent~~fluorescence was emitted from
the spot at three points in the tenth probe region, at two
points in the 41st probe region, and at one point in the 46th
probe region, respectively.

Spot positions in which fluorescence in association with
35 the formation of hybrid substances was observed in the prove

5 region having these base sequences including variations
corresponded to spot positions of weak fluorescence intensity
in the probe region having the above described 42nd original
base sequence. Thus, if the base sequences of the probes are
10 compared between both ~~the~~ regions, the base sequence of the
tenth probe is ACTCCG, the base sequence of the 41st probe is
the CCTCCA, and the base sequence of the 46th probe is CCTCTG
with respect to the original base sequence of CCTCCG ~~that of~~
the 42nd probe ~~has~~. For their complementary sequences, it can
be understood that with respect to ~~the~~ CGGAGG in the 42nd
15 probe, the CGGAGT and G were changed to T in the tenth probe,
the TGGAGG and C were changed to T in the 41st probe, and the
CAGAGG and G were changed to A in the 46th probe. That is, it
was confirmed that in test samples, forming hybrid substances
with these tenth, 41st and 46th probes, cDNA fragments
20 contained therein derived from the p53 gene caused one base
mismatch with respect to the 42nd probe due to the above described
variations.

By this method, existence/~~not-non~~-existence of variations
and types thereof could be detected at the same time for all
25 the 64 types of test samples.

(Example 6)

Preparation of a ~~substrate for probe matrices~~
~~partitioned by patterns~~. Substrate for Probe Matrices
Partitioned by Patterns

30 A glass substrate with an epoxy group introduced to the
surface and with black matrices for probe matrices was
prepared in accordance with the following procedure.

1. Introduction of an epoxy group to the surface of the
substrate-

35 A glass substrate made of synthetic quartz (50 mm x 50
mm) was first subjected to ultrasonic cleaning using a 2%

5 sodium hydrate solution, and was then subjected to UV ozone
processing to clean the surface. A 50% methanol solution
containing 1% of a silane coupling agent (trade name: KBM
403; manufactured by The Shin-Etsu Chemical Co., Ltd.)
containing a silane compound having an epoxy group bonded
10 thereto (γ -glycidoxypyrpyltrimethoxysilane) was stirred at
room temperature for three hours to perform a preliminary
treatment for hydrolyzing the methoxy group in the silane
compound. This solution already subjected to the hydrolysis
treatment was applied to the surface of the above -described
15 clean substrate with a spin coater, and was heated and dried
at 100°C for 5 minutes to form a binding coating of the silane
coupling agent on the surface of the substrate. Through the
formation of this coating, the epoxy group contained in the
silane compound was introduced to the surface of the
20 substrate.

2. Formation of black matrices-

Then, A DEEP-UV resist containing carbon black (negative
type resist for black matrices) (trade name: BK-739P;
manufactured by Nippon Steel Chemical Co., Ltd.) was applied
25 on the surface of the substrate with a spin coater, so that
the film thickness after curing was 5 μ m, and it was heated
for curing on a hotplate at 80°C for 5 minutes. By the
proximity exposure using a DEEP-UV aligner, a pattern was
exposed to light using as an exposure mask a mask for
30 negatives with patterning applied to a region of 10 mm x 10
mm, so that the distance X between adjacent wells was 100 μ m
and the outer shape of the well was a ~~square of~~ 1 mm x 1 mm
square. Then, the development was carried out with ~~a~~
~~developer of an~~ inorganic aqueous alkaline developer solution
35 using a spin drier, and the substrate was washed with pure

5 water to completely remove the developer~~-completely~~. Then, it
was briefly dried using the spin drier, and was thereafter
heated in a clean oven at 180°C for 30 minutes to fully cure
the resist. As a whole, a substrate with 400 wells ~~arranged~~
10 in a predetermined arrangement and black matrices (resist
walls) partitioning adjacent wells was obtained. Furthermore,
the internal volume of each well is calculated as 5 µl if the
thickness of the solution is 5 µm. Also, in the surface of
the prepared black matrix, the angle of contact to water was
93 degrees and wettability with water was significantly low,
15 while in the bottom of the well, the angle of contact to water
was 35 degrees and the wettability with water was high.

3. ~~Fixation of~~Fixing probe DNA-

64 oligonucleotides of 18 mer with an amino group bound
to the hydroxyl group of the five prime-end through a
20 phosphate group and hexamethylene were prepared as DNA probes.
The 64 probes are same as those prepared in Example 5 ~~as with~~
respect to base sequences, but are different in the sense that
an amino group is introduced in ~~its~~their five prime-end
instead of a thiol group.

25 5 µl of solution of these DNA probes was injected into
each well under a microscope, and was left in a humidified
chamber to allow the probe to bind to the substrate through
the reaction between the amino group of the five primer-end
and the epoxy group on the substrate.

30 ~~(Example 7)~~

Analysis of cDNA derived from the p 53 gene that has been
prepared from mRNA, using the probe matrix substrate
partitioned by the pattern that has been prepared in Example
6.6 is as follows.

5 As in the case of Example 4, 64 types of labeled cDNAs
were spotted in each probe region of 2 mm square as an
arrangement of 8 x 8 spots, as shown in FIG. 8, using a pin
system array preparing apparatus.

 A hybridization reaction was carried out by a method
10 similar to that ~~of example 4.~~ in Example 4.

 The obtained result was similar to that ~~of~~ in Example 5.

5 SEQUENCE LISTING

<110>Canon INC.

<120>An assay of many samples for multiple items at the same time

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10 <160>64

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40 <223>Sample ~~originucleotide~~oligonucleotide

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5 ABSTRACT OF THE DISCLOSURE

Multiple specimens, typically biological samples having different properties and origins, are bound onto matrix substrates, and oligonucleotides, proteins and drugs are spotted on each matrix in an array to examine those specimens
10 at ~~a~~the same time for multiple items.